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TITLE: SUICIDE INHIBITORS OF REVERSE TRANSCRIPTASE IN THE
THERAPY OF AIDS AND OTHER RETROVIRUSES

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INDEX

INDEX	PAGE
SUMMARY	1
A INTRODUCTION	3
B WORK ACCOMPLISHED	5
1. EPOXIDE SUICIDE INHIBITORS	5
2. SYNTHESIS OF URIDINE 2'3' RIBOSPIROXIRANES	6
3. ANTIVIRAL ACTIVITY OF URIDINE SPIROXIRANES AND CYTOTOXICITY SCREENING	8
4. ANTIVIRAL PROPERTIES OF STEPOL PHOSPHONOFORMATES	10
5. SYNTHESIS, ANTIVIRAL ACTIVITY, AND PHARMACOKINETIC RADIOLABELLED AZT-STEROI, DICARBOXYLATES	12
6. SYNTHESIS OF NUCLEOSIDE ANALOG TRIPHOSPHATES AND ENZYME KINETIC STUDIES	15
7. EXPRESSION OF HIV REVERSE TRANSCRIPTASE IN PFA SENSITIZING CELL LINES	20
APPENDIX	
I. PUBLICATIONS	24
II. COMPOUNDS SYNTHESIZED AND PREPARED FOR SHIPMENT TO USAMRIID FOR ANTIVIRAL TESTING	25
III. TEST DATA ON ANTI HIV DRUGS	27
IV. TEST DATA ON OTHER VIRUSES AND CONFIRMATORY SCREENING	39
V. SUMMARY TABLE OF ANTIVIRAL COMPOUNDS	77
VI. SUMMARY OF SIGNIFICANT PROJECT ACCOMPLISHMENTS RECOMMENDATIONS FOR FUTURE EXPLOITATION OF FINDINGS	78

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SUMMARY

This project was designed to develop antiviral agents by adapting new enzymatic and pharmacokinetic principles to inhibition of the HIV-reverse transcriptase. Towards this end synthetic organic procedures were directed to synthesis of compounds in the following categories.

1. 3' Nucleoside spiroxiranes and a series of related compounds having the potential to function as suicide inhibitor of the reverse transcriptase through the 3' functionality.
2. A series of sterol phosphonoformate analogs designed to improve delivery of the active phosphonoformate (PFA) moiety to sites of viral replication.
3. A series of 5' sterol ester derivatives of azido thymidine designed to improve the pharmacokinetics and blood half life of AZT.

Synthetic compounds were tested for antiviral activity against HIV and EIAV in tissue culture, against purified HIV-reverse transcriptase in enzyme kinetic studies, where appropriate and samples of synthetic analogs were supplied to the US Army Antiviral Program for screening against a battery of 10 viruses of interest as military disease hazards.

In addition to the targeted objectives, the serendipitous observation of a cell product that enhances the sensitivity of HIV-reverse transcriptase towards inhibition by Foscarnet (PFA) by almost 1000 fold, was explored further resulting in preliminary identification of the sensitizing compound.

The major accomplishments in the categories listed above are summarized below and are described in more detail in the appropriate sections of the final report.

Organic synthetic procedures for the synthesis of a new class of compounds, the 3' nucleoside spiroxiranes were developed. A series of these and related compounds and intermediates modified at the 2' and 3' functionalities were tested for activity against HIV and EIAV in tissue culture. Six compounds showing promising antiretroviral activity were identified. Several additional compounds were identified in the subsequent USAMRIID screening tests as being active against additional viruses including Punta, Toro, Vaccinia, and Yellow Fever.

A procedure for synthesis of sterol esters of phosphonoformate (FOSCARNET) was developed by

condensation of cholesterol chloroformate with the appropriate dialkylphosphite. Several of these compounds displayed greater antiviral activity than the parent compound PFA and side chains that enhanced (or detracted) from the activity were identified.

Three 5' cholesterol ester derivatives of AZT were synthesized and showed antiviral activity in tissue culture demonstrating that the cholesterol ester hydrolases of the cell could liberate the active moiety intracellularly. The ³H radiolabelled AZT derivative of cholesterol sebacate was synthesized and used to evaluate blood and tissue half life of the ³H-AZT moiety in mice. Administration of AZT as the sterol ester derivative improved the blood half life of AZT by over 100 fold (< 1/2 hr to > 48 hr) and was accompanied by significant accumulation in tissues including brain.

The interaction of the effective antiviral compound 3' uridine spiroxirane and the recombinant HIV reverse transcriptase enzyme was investigated in depth in order to characterize the suicide inhibition native of the interaction. This compound was designed as a potential suicide inhibitor of reverse transcriptase through the oxirane functionality at the 3' position. This compound showed good antiviral activity against HIV in cell culture and also against a second retrovirus (Equine Infectious Anemia).

In order to determine if the observed antiviral activity was due (as predicted) to inhibition of reverse transcriptase, the kinetic properties of 3' uridine spiroxirane were evaluated against the purified recombinant HIV-reverse transcriptase in vitro. In order to do this the triphosphate derivative of the nucleoside analog (the true putative inhibitor) was synthesized using a newly developed procedures for synthesizing nucleoside triphosphates.

It was found that the 3' uridine spiroxirane triphosphate derivative was an effective inhibitor of the HIV reverse transcriptase in the 0.1 to 1 micromolar range. The time course of the inhibition was progressive as expected for a suicide type inhibitor. Furthermore, the inhibition was not reversed by addition of excess template thus distinguishing it from a simple chain-terminating inhibitor and confirming that the inhibition resulted in the progressive and permanent inactivation of the enzyme characteristic of a suicide inhibitor. Thus it is concluded that the antiviral activity of the spiroxirane analogs we have synthesized is indeed related to their ability to function as suicide substrates for the HIV reverse transcriptase.

The compounds synthesized and supplied to USAMRID for antiviral testing are tabulated. The test data

indicating I_{50} , cytotoxicity and therapeutic index for all compounds showing antiviral activity are summarized on the Appendix to this final report.

The project was thus characterized by an unexpectedly high degree of success.

Of a total of 4a compounds synthesized and supplied to USAMRID for antiviral testing a total of 9 proved to have effective activity against one or more viruses of interest as military disease hazards including HIV, Punta, Toro, Vaccinia, and Yellow Fever. These compounds and the viruses against which they are effective are listed on the final page of this report. Furthermore, most of these showed low cytotoxicity in cell culture assays suggesting that they will be potentially useful therapeutic agents if future in vivo testing is ever carried out. Finally, the basic enzyme kinetic studies carried out in the extended (unfunded) 4th year of the contract, support the basic idea underlying these investigations, in that effective antiviral agents can be developed based upon the principle of suicide inhibition of key viral enzymes.

A. INTRODUCTION

Retroviruses are characterized by the anomalous storage of their genetic information in the form of RNA. They are a diverse group of organisms which have been shown to be causative agents in a number of mammalian and avian diseases states. Lentiviruses are a subfamily of retroviruses which have been linked to the induction of arthritis, encephalitis, and slow neurological diseases in certain species. Many of these viruses are characterized by their ability to develop novel antigenic variants that can escape temporarily from host immune surveillance. Interest in retroviruses in human disease states has increased with the isolation of HTLV I and II as the causative agents of adult human T cell leukemias and of HIV I and II as the cause of Acquired Immunodeficiency Syndrome (AIDS).

The most recent evaluation of the AIDS epidemic in the U.S. indicates that the current total of approximately 70,000 AIDS cases is accompanied by a further 1 to 1.5 million infected individuals, the majority of whom will progress to the clinically overt disease state in the next 5 years. In the near-term absence of an effective vaccine an increase in this infected/symptomatic category is likely to continue for a number of years.

In addition to an antiviral therapy for acute cases of AIDS therefore, there is clearly pressing need for an effective antiviral drug regimen suitable for administration to large numbers of otherwise healthy individuals in order to arrest progression of the disease. By preventing viral replication, such therapy may also reduce dissemination of the virus and thus indirectly fulfill some of the immediate functions of a vaccine. The current drug of choice, AZT can have undesirable side effects when used at maximum therapeutic dosages over prolonged periods. In addition HIV-variants with increased resistance to AZT have recently begun to appear. Combination therapy with drugs having a different mechanism of action can reduce the likelihood that resistant variants will appear. The drugs being developed in this project are designed with the above potential for use in long-term combination therapy low of toxicity as a primary goal.

Replication of retroviruses is critically dependent upon a single enzyme called reverse transcriptase (RT). This is an RNA-dependent DNA polymerase first found in the purified virion of Rous sarcoma virus by Temin and murine leukemia virus by Baltimore. The enzyme transcribes the viral RNA into DNA in the first step of viral replication. Since host cells do not contain reverse transcriptase, several molecules of the enzyme are packaged in each virion and enter the cell together with the viral RNA. It is at this stage that viral replication is most sensitive to inhibitors of the reverse transcriptase, before amplification of the viral genome and pol gene directed synthesis of endogenous RT has taken place.

The principle that inhibitors of reverse transcriptase will inhibit replication of retroviruses is well established. For example, 3'-azido-3'-deoxythymidine, the triphosphate of which inhibits the RT activity of HIV, is a potent inhibitor of virus replication in cultured H-9 cells in the range of 1-10 micromolar and is being used successfully in patients with AIDS. Prolonged administration may cause serious side effects. Another agent which has shown promise is phosphonoformate (PFA Foscarnet) which inhibits the RT activity of HIV with an I_{50} of only 0.1 micromolar. However, much higher concentrations of up to 340 micromolar are required for complete inhibition of HIV replication in H-9 cells. Other drugs including the dideoxycytidines which are also based upon inhibition of RT by chain termination of the viral template are in clinical trial. Since none of these drugs permanently inactivate the reverse transcriptase and since they do not accumulate intracellularly in significant amounts, virus replication will resume when blood levels of the drugs decrease.

Since there is no complete animal model for HIV which reproduces the immunodeficiency, as an adjunct to the tissue culture studies this project will also utilize the Rauscher murine leukemia virus which has a reverse transcriptase with close homology to that of HIV, for *in vivo* testing of new antiviral drugs. This will be supplemented by testing drug toxicity in human T lymphocytes growing in tissue culture.

Suicide Inhibitors of Reverse Transcriptase

Suicide inhibitors, also known as Kcat inhibitors, are active site directed substrate analogs which contain a latent, reactive group. Following cleavage of the analog by the target enzyme the reactive group is released at the active site and inactivates the enzyme. Inhibitors of this type have several potential therapeutic advantages in that they can be designed to be highly specific for

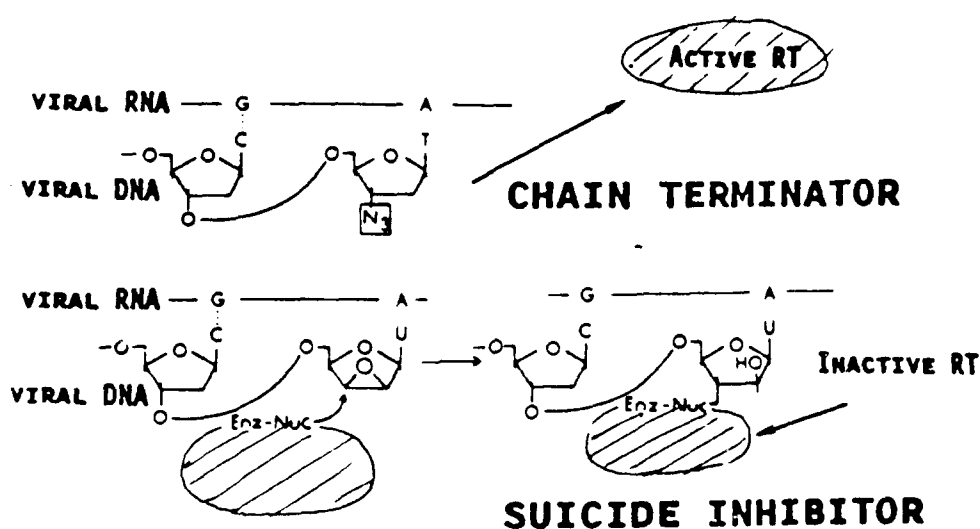
the target enzyme. Since most normal cells do not contain active or functional reverse transcriptases this reduces the possibility of side effects. Furthermore, by their nature, inhibitors of this type remain metabolically inert unless specifically cleaved by the target enzyme. Finally, the inhibition is permanent and irreversible, so that viral replication will not resume when the blood levels of the drug decrease. This feature could be particularly useful in long-term therapy of chronic human retrovirus infection.

Inhibition of retroviral replication by 3'-Nucleoside Spiroxirane Derivatives:

An apparent suicide inhibitor of E.Coli DNA polymerase was described by Abboud et al. It was shown that adenosine 2'3'-riboepoxide 5'-triphosphate irreversibly inactivated the enzyme by a covalent interaction. Because of the intrinsic reactivity of the epoxides however they are not suited for use under physiological conditions. We will therefore synthesize a series of nucleoside 3'-spiroxirane analogs. These are designed to inhibit the enzyme by an analogous suicide mechanism and are based upon the observation that the uridine 3'-ribospiroxirane derivative is an inhibitor of retroviral replication in cultured cells and selectively suppresses reverse transcriptase production by the VCF 21 (Moss) vaccinia HIV-RT recombinant. Furthermore at therapeutic levels it has no significant effects on ^3H -thymidine incorporation in cultured T-lymphocytes.

In the first step of replication of viral DNA by reverse transcriptase, activation of the primer 3'-OH by the enzyme results in incorporation of the suicide nucleotide by a normal 3',5'-phosphodiester linkage. At this point the inhibition resembles a simple chain termination similar to that produced by other 3'-blocked nucleoside analogs such as 3'-azidothymidine (AZT) and dideoxycytidine. However when the next step in the elongation reaction is attempted, the enzyme by activating the 3'-oxygen of the spiroxirane functionality becomes alkylated by the substrate and effectively commits suicide. A primary objective of these studies will be to characterize the kinetics of inhibition of the purified HIV-reverse transcriptase by the nucleoside spiroxirane triphosphates to determine if this second suicide step occurs and contributes to the observed antiviral activity of these compounds. Initial indications that the analogs can accomplish irreversible inactivation of intracellular RT support this possibility, and further efforts will be made to incorporate features into the molecule which favor the suicide pathway,

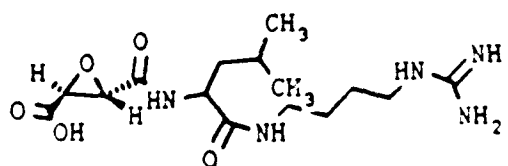
Because of the greater stability of the oxirane ring under physiological conditions these analogs offer several advantages over the simple epoxide types: A schematic representation of the consequences of an interaction with a suicide inhibitor versus a chain terminator is given below.



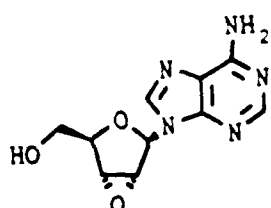
B. WORK ACCOMPLISHED

Epoxide Suicide Inhibitors:

Epoxide-containing irreversible (suicide) inhibitors have been reported for a number of enzymes, the structures of two such compounds are shown below. One of the first, [N-(L-3-*trans*-carboxyloxiran-2-carbonyl)-L-leucyl]-amido (4-guanido) butane was isolated from *Aspergillus japonicus* and was found to irreversibly inhibit cysteine proteases by alkylation of the sulphydryl group at the active site.

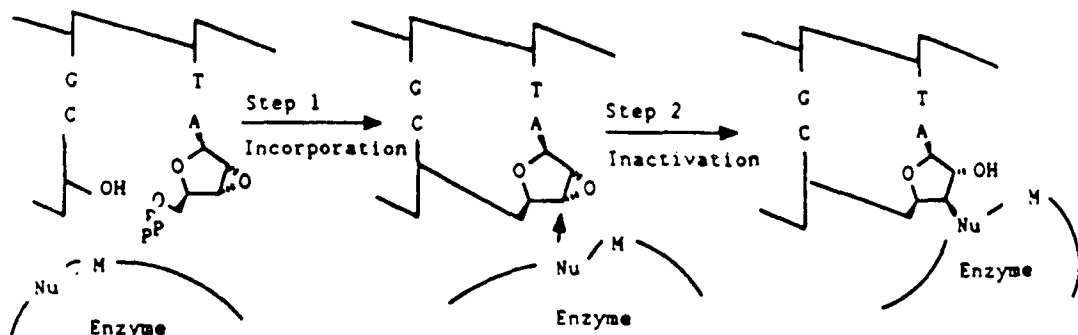


[N-(L-3-*trans*-carboxyloxiran-2-carbonyl)-L-leucyl]-amido(4-guanido)butane



9-(2,3-Anhydro- β -D-ribofuranosyl)adenine

A 2',3'-riboepoxynucleoside, 9-(2,3-anhydro- β -D-ribofuranosyl)adenine, was shown, as the 5' triphosphate, to irreversibly inhibit avian myeloblastosis virus DNA polymerase (a reverse transcriptase). Its mechanism of inactivation, shown below, is believed to involve alkylation of the enzyme in the active site upon activation of the epoxide ring after incorporation of the nucleoside to the end of the growing DNA strand.

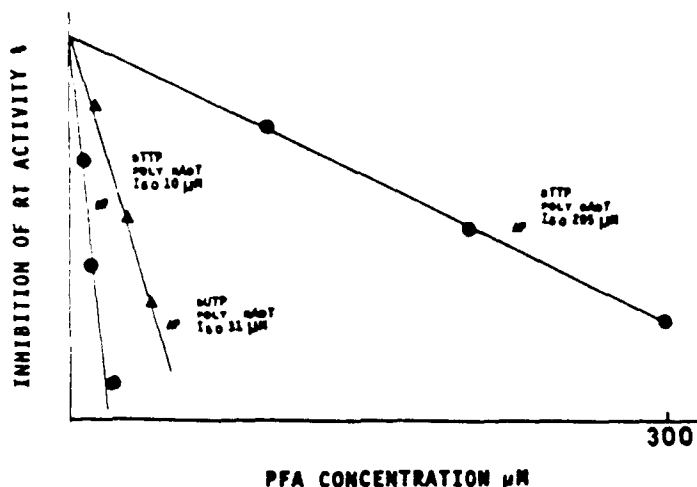


A proposed mechanism for RT inactivation by 2',3'-riboepoxyadenosine.

A similar compound, 1-(2,3-anhydro- β -D-lyxofuranosyl)cytidine, was recently synthesized by Broder et. al. and was found to inhibit HIV *in vitro*. Although the mechanism of inactivation was not elucidated, irreversible inactivation was considered a possibility. We have also synthesized the corresponding 1-(2,3-anhydro- β -D-lyxofuranosyl) uridine and found it to have antiviral activity. However these epoxide derivatives unlike the spirooxiranes (see below) are also cytotoxic at higher concentrations probably owing to the greater reactivity of the epoxide ring.

Preliminary Synthetic and Antiviral Screening Studies:

The reverse transcriptase of HIV is a highly error prone enzyme, a factor which probably contributes to the high mutation rate of the HIV genome. We have shown that reverse transcriptase readily incorporates dUTP in place of dTTP when transcribing from a poly rAdT template, and this incorporation, like that of dTTP, is particularly sensitive to inhibition by PFA (figure).



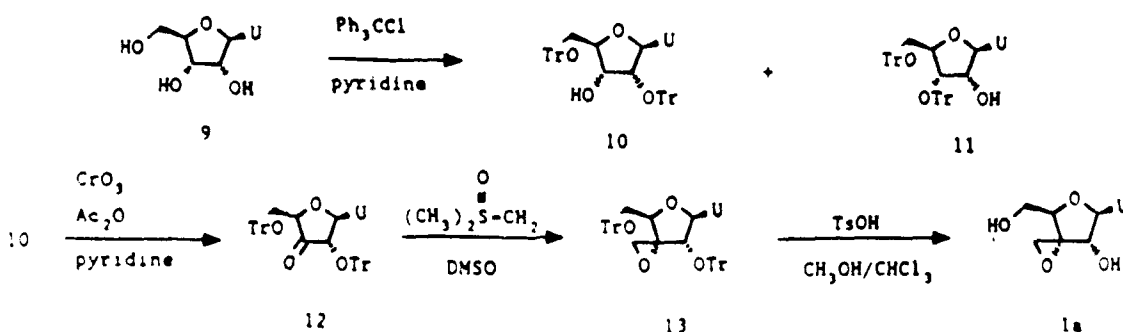
Reverse Transcriptase catalyzed Incorporation of d-UTP: Differential Inhibition of First-strand Synthesis by Foscarnet.

Since uridine is an unnatural base for the cellular DNA polymerases, initial studies were carried out using uridine nucleosides to enhance the antiviral selectivity.

Synthesis of Uridine 2' and 3'-Ribospiroxiranes:

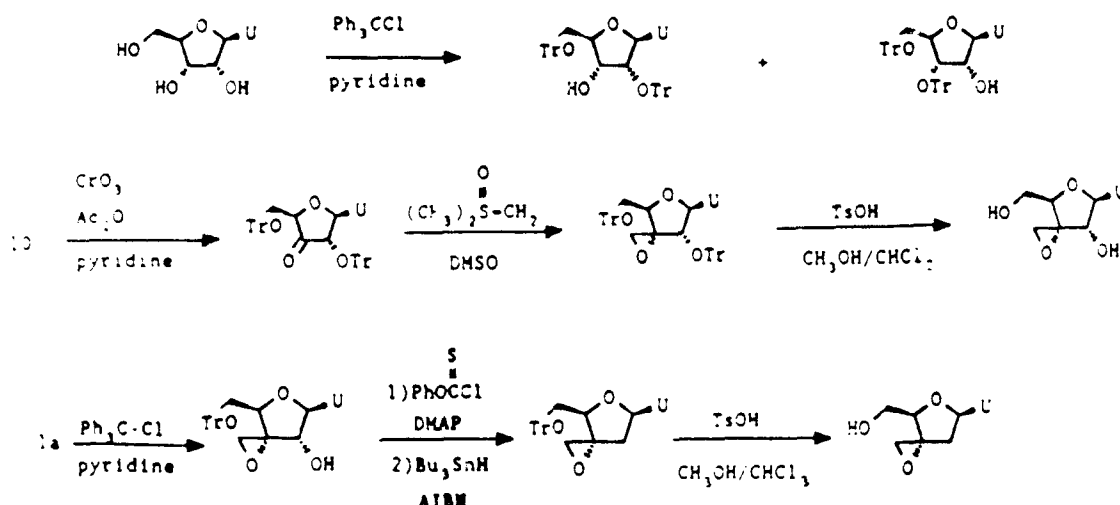
3'-uridine spiroxirane was synthesized from uridine by the sequence of reactions shown below. The procedure begins with the selective tritylation of uridine by the procedure of Cook and Moffat (38). Chromatographic separation of the isomers yielded the 2'S' and 3'S'-di-O-trityluridines in good yield. The 2',5'-isomer was oxidized with chromium trioxide in pyridine/acetic anhydride to 3',5'-di-O-trityl 3'-ketouridine. Treatment of this compound with two equivalents of dimethyloxosulfonium methylide yielded the 2',5'-di-O-trityl spiroxirane derivative which upon mild acid catalyzed hydrolysis yielded the deprotected 3'-uridine spiroxirane.

The corresponding 2'-spiroxirane derivative was also prepared from the 2',3'-di-O-trityl isomer by analogous procedures and characterized by IR and NMR spectroscopy.



Synthesis of Spiroxirane Nucleosides:

This will be carried out by the general procedures used for synthesis of the uridine 3'-ibospiroxirane described in section C above. The further extension of this to the synthesis of the 2'-spiroxiranes from the 3'5'-di-O-trityl isomers and the 3'-dideoxy spiroxiranes from the 5'-mono-O-trityl derivative is summarized in the scheme below.



This route to the deoxynucleoside spiroxiranes is preferred since direct conversion of the corresponding deoxynucleosides by procedures analogous to steps 1-3 in the above scheme may lead to base elimination following conversion to the 3'-keto derivative. Since trityl ethers of primary alcohols are more readily hydrolyzed than those of secondary alcohols it is not possible to obtain the 5'-mono-trityl uridine spiroxirane directly from the 3',5' derivative, which must first be completely eprotected and converted by the procedure of Michelson and Todd. Free radical deoxygenation of C-2 by the procedure of Acton et al via the 2'-O-phenylthiocarbonate followed by eprotection of the 5'-hydroxyl group yields the desired 2'-deoxy-3'-spiroxirane. The absolute configurations of the 3' and 2' uridine spiroxiranes obtained by this synthesis have not yet been assigned. However it has been reported by Corey and Chaykowsky that dimethyloxosulfonium methylide affords spiroepoxides with an equatorial methylene in reaction with 4-*t*-butyl cyclohexanone, which would lead to spiroxiranes having the ribo-configurations illustrated. Spiroxiranes in the alternative lyxo configuration however may also be obtained by this general procedure by reaction of the 3'-ketoderivatives with dimethyl sulfonium methylide, as illustrated in the scheme below. In contrast to dimethyloxosulfonium methylide, this reagent yields spiroexpoxides with an axially oriented methylene

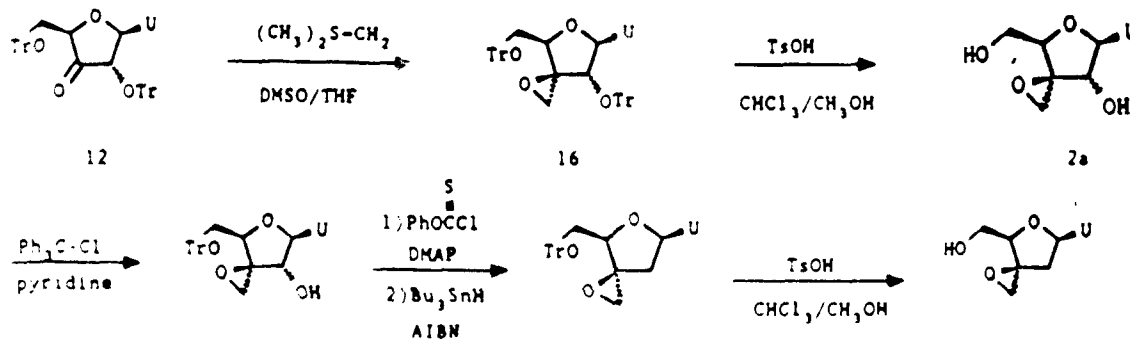
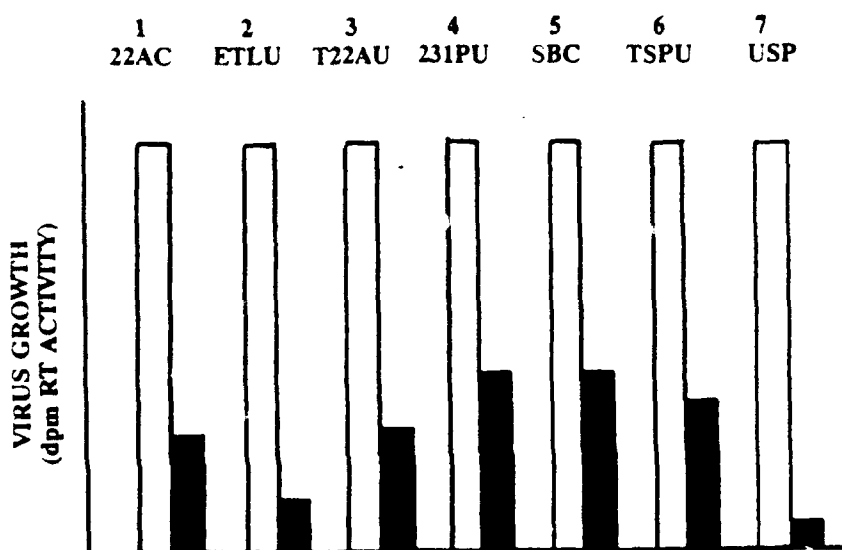


Fig.

Antiviral Activity of Uridine Spiroxiranes and Cytotoxicity Screening in Human T-Lymphocytes:

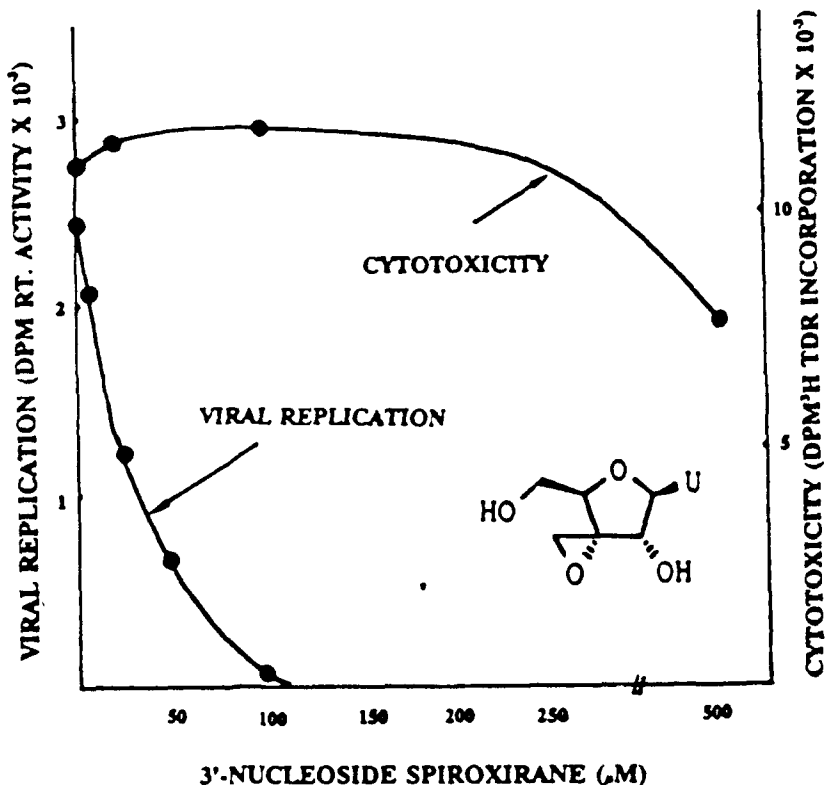
A series of approximately 30 nucleoside analogs, suggested by the considerations outlined in section B, were synthesized. These were screened for ability to inhibit replication of equine infectious anemia virus growing in equine dermal fibroblasts. This is a lentivirus having a close homology with HIV and is non pathogenic to humans. It is used as a preliminary screen for antiviral drugs before testing against HIV grown in A 3.01 human T-lymphocytes as described in section D. Cytotoxicity was evaluated at concentrations from 25-500 μ M in IL-2 supplemented CTLL T-Lymphocyte cultures pulsed with 3 H thymidine at 24 and 48 hours.



Antiviral Activity of Some Epoxy- and Oxirane Nucleosides:

Viral replication was monitored by RT assay on the pelletized virions from the culture medium. As illustrated in the figure above, 7 of these compounds inhibited viral replication by greater than 50% when present in the culture medium at a concentration of 100 μ M. The two most effective analogs were the uridine 2', 3'-lyxo epoxide (compound 2) and the uridine 3'-ribospiroxirane (compound 7). Both compounds were also tested for cytotoxicity by measuring their effects on 3 H-thymidine incorporation in the IL-2 dependent CTLL T-lymphocyte cell line in tissue culture.

The 3'-uridine spiroxirane derivative was added to control and EIAV infected cultures over a range of concentrations from 5 to 100 μ M and reverse transcriptase activity was assayed on the pelletized virions after 8 days. Virus replication was almost completely inhibited by 100 μ M concentrations of the 3' uridine spiroxirane the I_{50} being about 25 μ M. The corresponding 2'-spiroxirane used as a control had no effect at similar concentrations. The 3'-spiroxirane was non-toxic to T-lymphocytes, the I_{50} for inhibiting 3 H-thymidine incorporation being greater than 500 μ M. Since the epoxide derivative was somewhat inhibitory to T-cell proliferation at concentrations above 100 μ M, whereas the spiroxirane analog was not, the nucleoside spiroxirane family has been selected for further in depth characterization in this project.



Antiviral Activity of Uridine 3'-ribospiroxirane.

Cultures of equine dermal cells were infected with a standard inoculum of the EIAV retrovirus in the presence of increasing concentrations of uridine 3'-spiroxirane. Virus replication was measured by following release of reverse transcriptase into the culture medium. In separate experiments growth inhibition of cultured T lymphocytes was evaluated by measuring incorporation of tritiated thymidine (TdR) into cellular DNA. Note the greater sensitivity of the viral versus the cellular polymerases presumably due to the ability of the cellular polymerases to "proofread" and excise the unusual nucleoside.

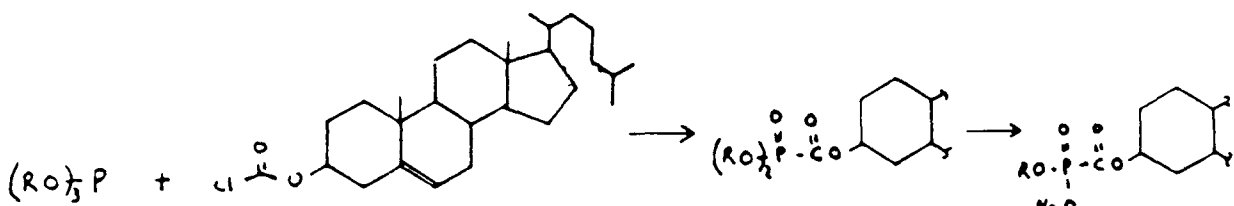
Antiviral Properties of the Sterol Phosphonoformates

Despite the sensitivity of the HIV-RT to inhibition by PFA, replication of the virus in tissue culture is relatively insensitive, the I_{50} being of the order of 50 μM .

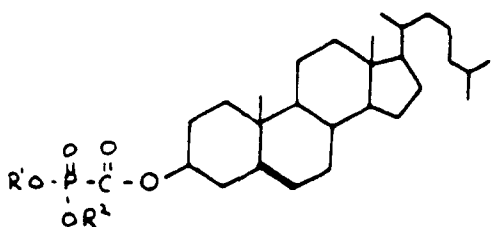
We have shown in previous studies, by use of sterol ester analogs such as the sterically hindered hydrolysis resistant cholesterol $\alpha\alpha$ -methyl ethyl caproate (CMEC), that intact cholesterol esters enter cells via a specific endocytotic transport process. To enhance deliverability of the active PFA moiety to endosomal sites of viral replication, cholesteryl phosphonoformate ester analogs of PFA have been synthesized and characterized.

Cholesteryloxycarbonylphosphonoformates:

A series of mono- and di-alkyl cholesteryloxycarbonylphosphonoformates have been successfully prepared via an Arbozov reaction between trialkylphosphites and cholesterylchloroformate.



Triesters gave acceptable spectroscopic characteristics and were selectively hydrolyzed to the mon and di-sodium salts



Properties of Some Sterol Phosphonoformates.

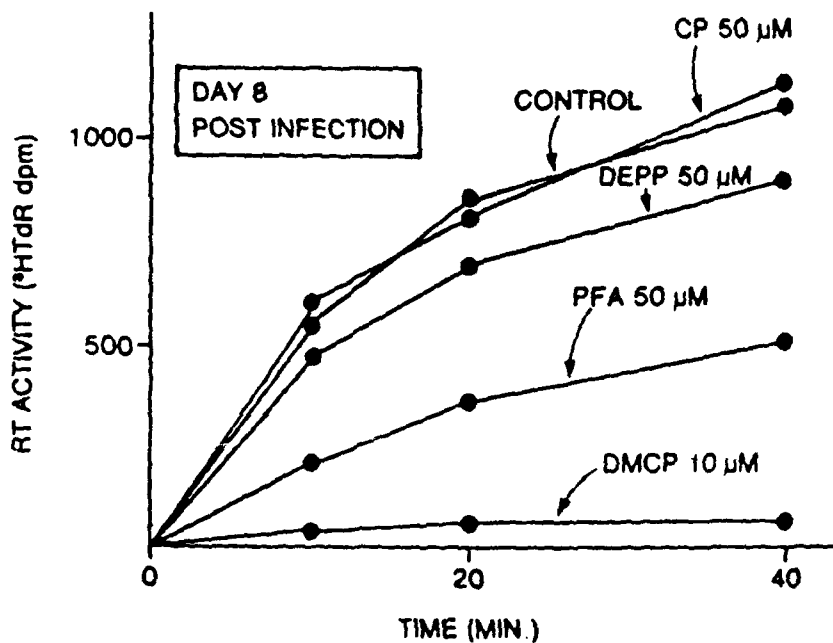
Cmpd,5	R ¹	R ²	MP (C°)	R _t
a	C ₂ H ₅	C ₂ H ₅	110-111	0.60 (A) ^o
b	CH ₂ CH ₂ CH (CH ₃) ₂	CH ₂ CH ₂ CH (CH ₃) ₂	67-69	0.51 (B)
c	CH ₂ CHO (CH ₃) ₂ OCH ₂	CH ₂ CHO (CH ₃) OCH ₂	106-108	0.30 (A)
d	CH ₂ CH ₂ CH ₂ CO ₂ C ₂ H ₅	CH ₂ CH ₂ CH ₂ CO ₂ C ₂ H ₅	oil	0.33 (A)
e	C ₂ H ₅	Na	≥220	0.29 (C)
f	Na	Na	≥300	-

^a Solvent systems: A, 50% ethyl acetate/hexanes, B, 20% ethyl acetate/hexanes, C, 52/14/1/1 CHCl₃/CH₃OH/con NH₄OH/H₂O.

A number of compounds of this type having both aromatic and aliphatic substituents have been tested for antiviral activity in vitro, as detailed below. Incorporation of the sterol moiety in an appropriate triest environment can result in compounds having 20-30 times the antiviral potency of the parent compound PFA. Table I lists the effect of increasing the number and size of the R groups on the melting point and chromatographic mobilities of some of these derivatives.

Pending development of the low biohazard CPE assay for HIV infectivity described below, the antiviral potency of the drugs was evaluated against Equine Infectious Anemia Virus (EIAV) growing in fetal equine fibroblasts. This is a lentivirus with a reverse transcriptase having the highest degree of homology with HIV-1 RT. The assay consists in measurements of RT activity on pelletized supernatants of infected cultures 8 days following infection. These sterol esters of phosphonoformate show an interesting pattern of antiviral activity in this system.

ANTI-VIRAL ACTIVITY OF SOME STEROL PHOSPHONOFORMATES



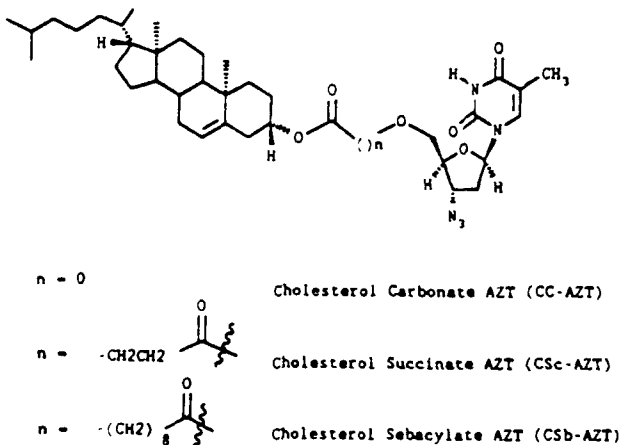
The approximate I_{50} for inhibition of EIAV replication by PFA is between 40 and 50 μM . The diethyl-phenyl-phosphonoformate ester (DEPP) is only about 20% as active as PFA and the doubly charged lithium cholesterol phosphonoformate (CP) is essentially inactive. Substitution of the phenyl group with a cholesterol group, however, results in a major enhancement of activity. For example, the closely related DMCP (lower curve, \square) in which cholesterol is esterified to the formate moiety, has 10-20 times more anti-viral activity than the parent compound PFA. Some of the more active analogs display activities close to the theoretical maximum predicted by the enzyme kinetic studies.

The sterol phosphonoformates display another useful property since the antiviral effects persist for a considerable time (up to 8 days) after removal of the drug from the culture medium. It seems likely that this prolonged antiviral protection may be due to intracellular accumulation of the analog followed by slow hydrolysis and sustained release of the active PFA moiety. One of the objectives in this Project is to explore this observation in more depth by measuring rates of intracellular accumulation and hydrolysis of sterol phosphonoformates using ^{14}C -labelled compounds synthesized by analogous procedures. In this way ligands which enhance uptake can be distinguished from those which enhance (or reduce) hydrolysis. This information will be used to enhance pharmacokinetic properties which may prove clinically useful.

SYNTHESIS AND ANTIVIRAL ACTIVITY OF AZT-STEROL DICARBOXYLATES:

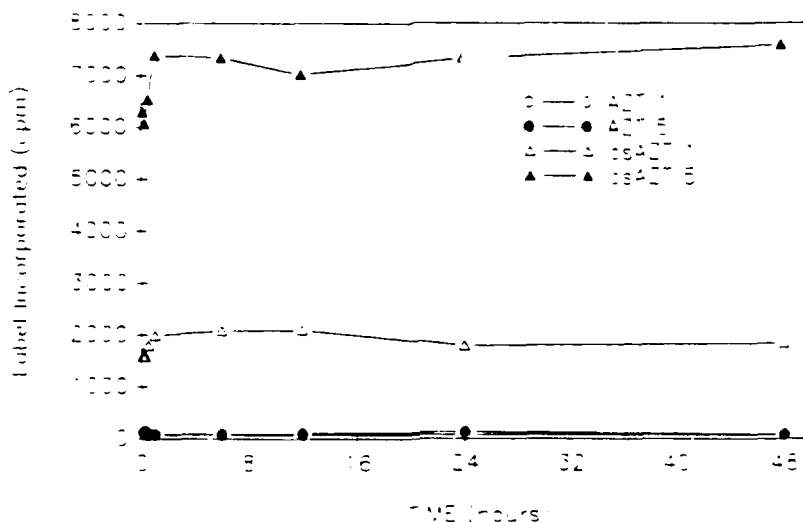
^3H -labelled AZT was prepared by NaB^3H_4 reduction of the 5'-aldehyde derivative and coupled cholesterol by a C_{10} (Sebacate) linker as described in section D, to yield cholestryl sebacyl ^3H -AZT (CS-AZT). Unlabelled cholestryl succinyl AZT (CS-AZT) and cholestryl carbonate AZT (CC-AZT) having the structures illustrated below, were also synthesized

STRUCTURES OF SOME SYNTHETIC AZT STEROL DIESTERS



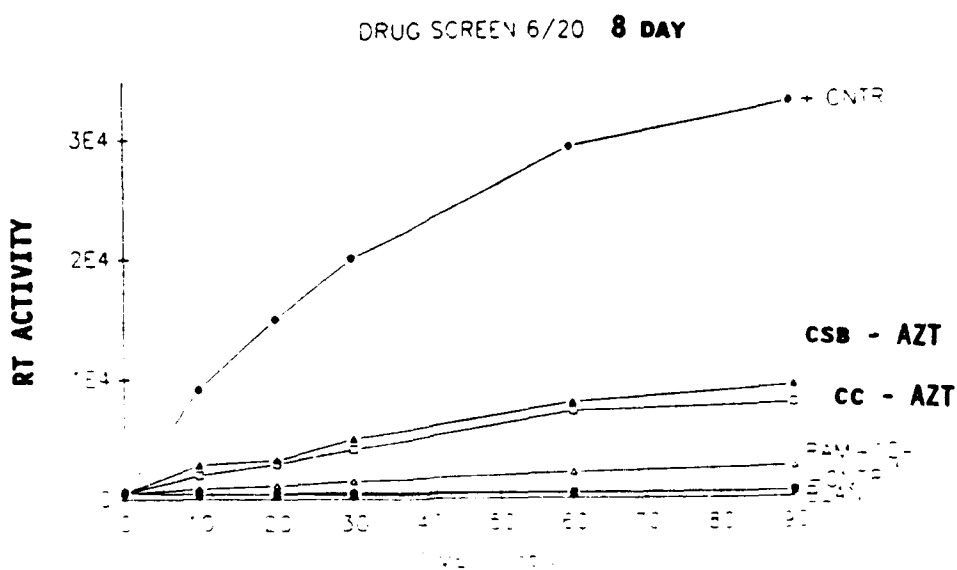
The uptake and accumulation of the ^3H -labelled AZT sterol sebacylate was compared with that of ^3H -AZT in the human T-lymphocyte cell line over a 48 hr period at two concentrations ($1 \mu\text{M}$ and $5 \mu\text{M}$) in the medium. Cells were harvested at intervals and washed using a Mash harvester before counting. Maximum accumulation and retention (following the wash procedure) occurred within 1 hour and was approximately 15 and 80 fold greater for the sterol derivatives than for free AZT at the two concentrations ($1 \mu\text{M}$ and $5 \mu\text{M}$) respectively.

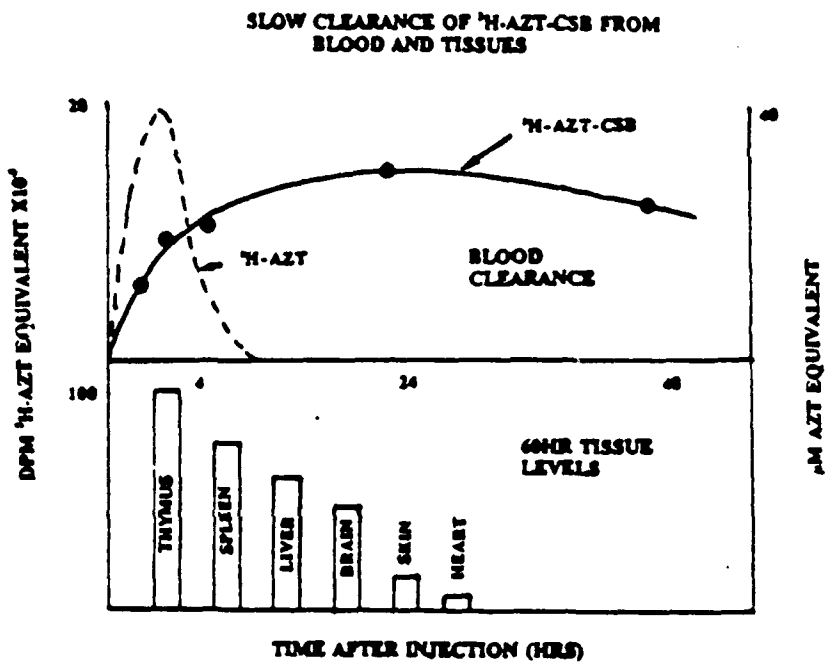
**ENHANCED ACCUMULATION OF RADIOLABELLED
CSB - AZT IN T LYMPHOCYTES**



The unlabelled AZT-sterol dicarboxylates showed good antiviral activity at similar concentrations as measured by suppression of viral RT levels in culture supernatants Figure suggesting that the products were being hydrolyzed intracellularly to the free form. Use of the radiolabelled analogs to explore factors regulating this intracellular accumulation and release will be a major objective of the further studies proposed in this project.

**ANTIVIRAL ACTIVITY OF
SOME STEROL AZT DIESTERS**





Tissue Distribution of ^3H -AZT-CSB 25 Hours and 60 Hours Following Injection of a Single 20mg/kg

Equivalent Dose

Tissue	25 Hrs			60 Hrs		
	Sample dpm	dpm per mg	$\mu\text{M AZT}$ equivalent	Sample dpm	dpm per mg	$\mu\text{M AZT}$ equivalent
liver	42596	38.4	33	57594	62.6	54
spleen	10247	183	159	9873	80.3	70
thymus	1606	61.8	54	1661	139	121
ovary	--	--	--	2273	59.8	52
skin	1398	20.9	18	1589	8.4	7
stomach	11500	134	117	10779	38.2	33
lung	2311	25.1	22	11324	104	90
heart	21275	154	134	707	4.5	4
kidney	18073	38.8	34	7183	37.8	33
brain	3914	9.3	8	3922	42.5	37
muscle	2043	16.7	15	4516	36.7	32

SYNTHESIS OF NUCLEOSIDE TRIPHOSPHATE ANALOGS FOR ANTIVIRAL TESTING:

The classical procedures for synthesis of nucleotide triphosphates require relatively large quantities of the nucleoside precursor and frequently give poor yields.

Ludwig and Eckstein (J. Org. Chem. 1989, 54, 631-635) have reported an approach involving condensation of an activated nucleoside derivative with inorganic pyrophosphate. In this synthesis 2-chloro-4H-1,3,20-benzodioxaphosphorin-4-one phosphorylates the 5'-hydroxy group of a nucleoside to form an intermediate 2, which on subsequent reaction with pyrophosphate produces a nucleosidylcyclotriphosphite 3. This intermediate is oxidized with iodine/water to furnish nucleoside 5'-triphosphate. The procedure is suitable for use with low milligram quantities.

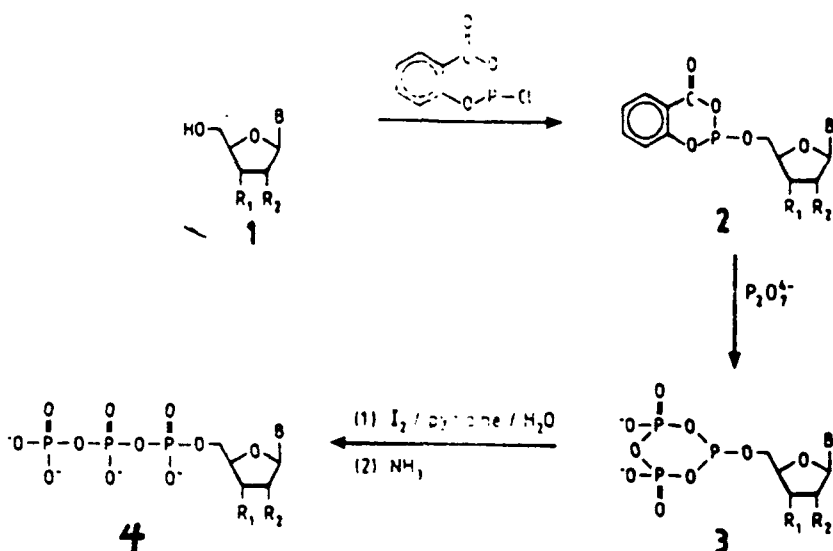
The series of reactions illustrated in Scheme I was successfully applied to microscale preparations of the 5'-triphosphates of both AZT and 3' uridine spiroxirane 1a which was previously synthesized and shown to have antiviral activity.

The formation of intermediate 3 was detected by ^{31}P NMR spectroscopy. The observed chemical shifts are comparable to those reported in the literature. The spectrum is of the ABX type where the AB part corresponds to the two phosphate groups with close but not identical shifts and the X part corresponds to the trivalent phosphorous atom. Oxidation of the intermediate 3 yields the normal triphosphate. In the ^{31}P NMR spectrum, the triplet of the trivalent phosphorous atom of the intermediate compound 3 disappears indicating complete oxidation.

We suspected some nucleotide by-products due to the contamination by a small amount of triphosphate. Also hydrolysis of intermediate 2 could result in phosphorous containing contamination detected by ^{31}P NMR. These by-products must copurify with the unreacted nucleoside on DEAE-cellulose chromatography.

The antiviral activity of the synthesized AZTTP was tested and proved comparable to, and in some cases even more effective than, a sample provided by the Burroughs Wellcome Co., thus validating the procedure. The nucleoside spiroxirane triphosphate may exist in two different isomeric forms which may or may not have identical antiviral activity. We have isolated one isomer that has proved antiviral activity. Isolation and characterization of the other isomer requires further investigation. The validated procedure was then applied to synthesize the 5' triphosphate of 3' uridine spiroxirane.

Scheme I



The phosphorylating agent 2-chloro-4H-1,3,2-benzodioxaphosphorin-4-one was purchased from Aldrich. Dimethylformamide (DMF) was dried over $MgSO_4$ and distilled under reduced pressure. Anhydrous dioxane was distilled from LAH. Anhydrous pyridine was prepared by fractional distillation of refluxing pyridine with potassium hydroxide.

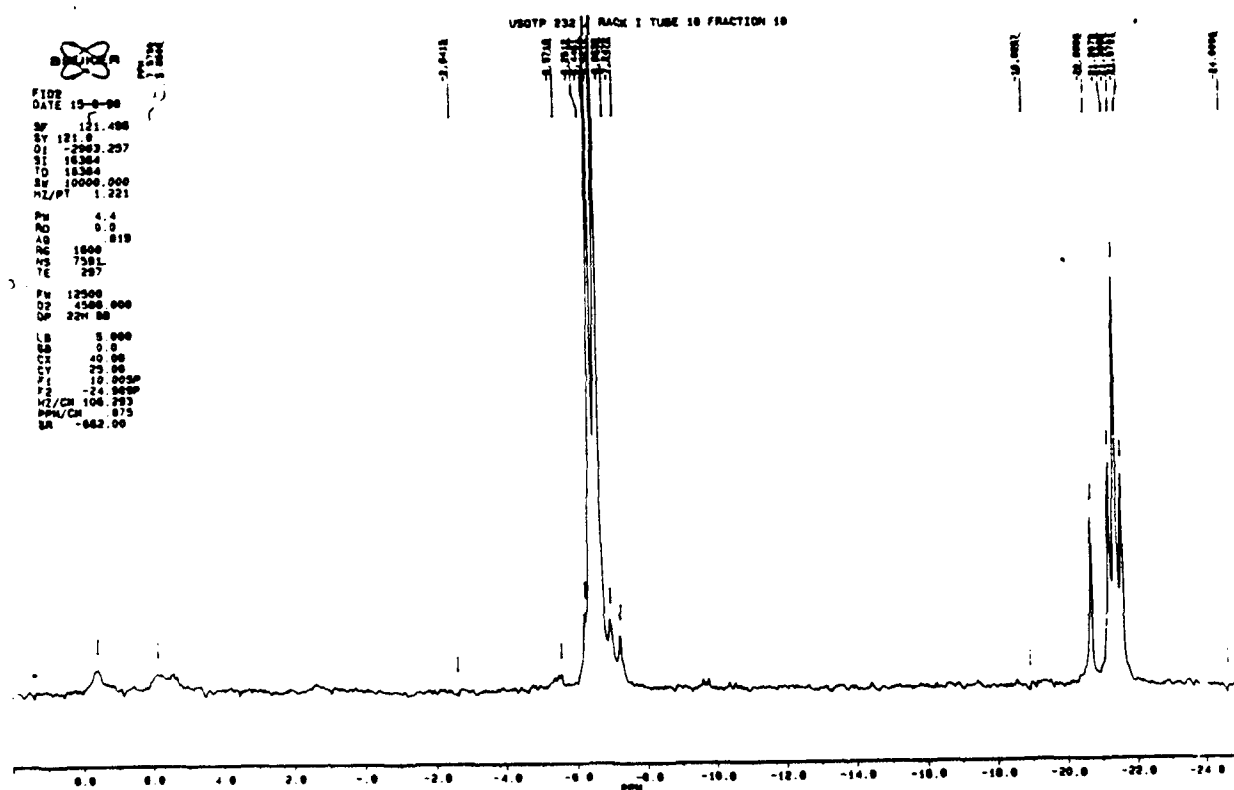
^{31}P NMR spectra were recorded on a Brucker 300 spectrometer with broad band decoupling. TLC was performed on either silica GHLF (Analtech) developed with isopropanol, ammonia, water (3:1:1) (system 1) or on DEAE-cellulose (Analtech) developed with 0.02 N hydrochloric acid (system 2).

Bis(tri-n-butylammonium) pyrophosphate. Tetrasodium diphosphate decahydrate (2.23 g, 5 mmol) was dissolved in water (50 ml), the solution was applied to a column of Dowex 50WXB in the H^+ form and the column was washed with water. The eluate was directly dropped into a cooled (ice water) and stirred solution of tri-n-butylamine (2.38 ml, 10 mmol) in ethanol (20 ml). The column was washed until the pH of the eluate increased to 5.0 (approximately 70 ml of water). The ethanol/water solution was evaporated to dryness and reevaporated twice with ethanol and finally with anhydrous DMF and diluted to 10 ml. This solution was stored over 4-Å molecular sieves.

Nucleoside spiroxirane 1a (100 μ mol) was dissolved in anhydrous pyridine/DMF, 1/4, V/V and evaporated to dryness in vacuo. The residue was dried further over P_2O_5 under reduced pressure for 2 hours at room temperature. The reaction flask was filled with nitrogen, during all the following manipulations a small positive pressure of nitrogen was maintained in the reaction vessel. Anhydrous pyridine (200 μ l) and DMF (800

μl) were injected through septum. A freshly prepared 1 M solution of 2-chloro-4H-1,3,2-benzodioxaphosphorin-4-one in anhydrous dioxane ($110 \mu\text{l}$, $110 \mu\text{l}$) was then injected into the well-stirred solution of nucleoside. After 15 minutes a well-vortexed mixture of a 0.5 M solution of bis (tri-n-butylammonium) pyrophosphate in anhydrous DMF ($300 \mu\text{l}$) and tri-n-butylamine ($100 \mu\text{l}$) was quickly injected and the reaction mixture was stirred for 10 minutes. A solution of 1% iodine in pyridine/water (98/2, V/V) (2 ml , $157 \mu\text{mol}$) was then added. After 15 minutes excess iodine was destroyed by adding a few drops of a 5% aqueous solution of NaHSO_3 and the reaction solution was evaporated to dryness. The residue was dissolved in water and applied to a DEAE-cellulose column which was eluted with a linear gradient of 800 ml of each 0.05 M and 1 M TEAB. The fractions were characterized by NMR spectroscopy. The presence of the characteristic triphosphate group was confirmed by NMR as indicated in the figure below.

NMR SPECTRUM OF 3' URIDINE SPIROXIRANE TRIPHOSPHATE DERIVATIVE



ENZYME INHIBITION STUDIES WITH 3' URIDINE SPIROXIRANE TRIPHOSPHATE

The recombinant HIV-reverse transcriptase was expressed in the vaccinia virus construct VCF21 as described in previous progress reports. The enzyme was purified from culture fluids by column chromatography on DEAE cellulose and carboxy-methyl cellulose columns. The purified enzyme was used to synthesize DNA using a poly rAdT template and following the enzyme activity by measuring incorporation of ^3H labelled dTTP. The synthesized ^3H DNA was collected on acid washed filter, and counted in a scintillation counter. It was found that 3' uridine spiroxirane was an excellent inhibitor of the enzyme in this system. The inhibition was time and concentration dependent consistent with the irreversible inhibition associated with a suicide inhibitor. More detailed kinetic studies on reversibility however will be required to establish this and to distinguish the kinetics observed from those of a chain terminating inhibitor.

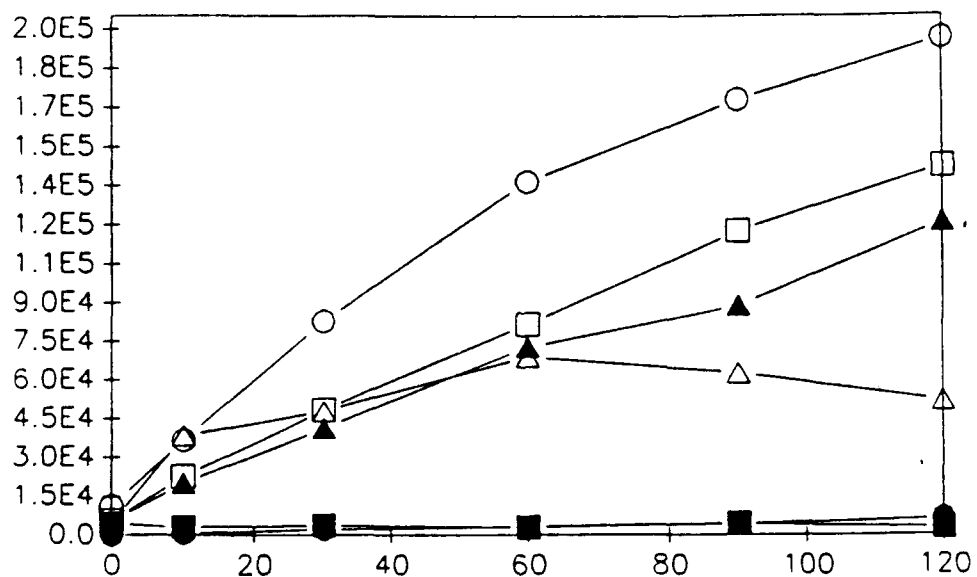


Figure: Activity of HIV-reverse transcriptase and inhibition by synthetic 3' uridine spiroxirane triphosphate

Key O control □ 0.05 μM ▲ .25 μM △ .25 μM ■ 2.5 μM ● Reagent blank

The triphosphate of 3' uridine spiroxirane was more potent as an inhibitor of the HIV reverse transcriptase than the triphosphate of AZT synthesized and tested in this system under the same conditions.

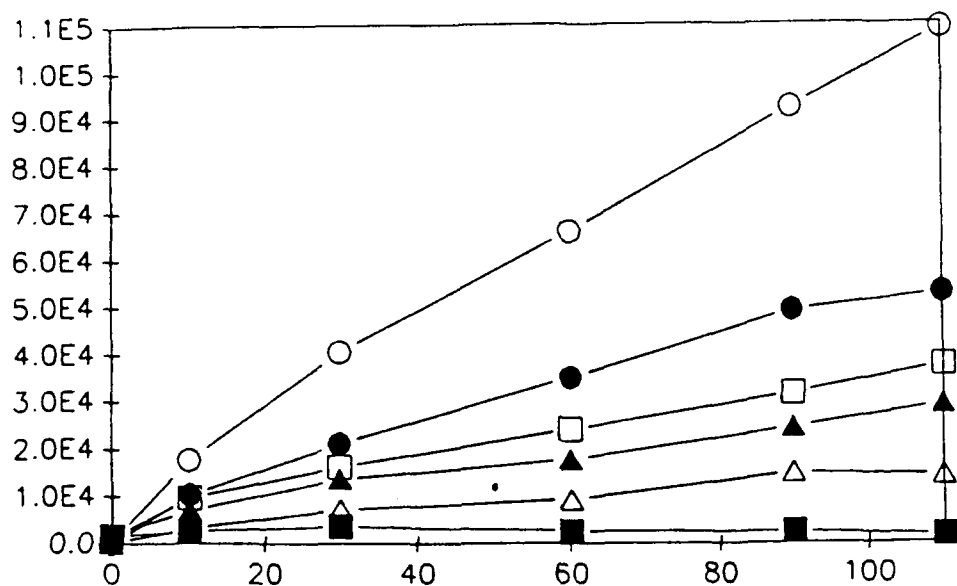


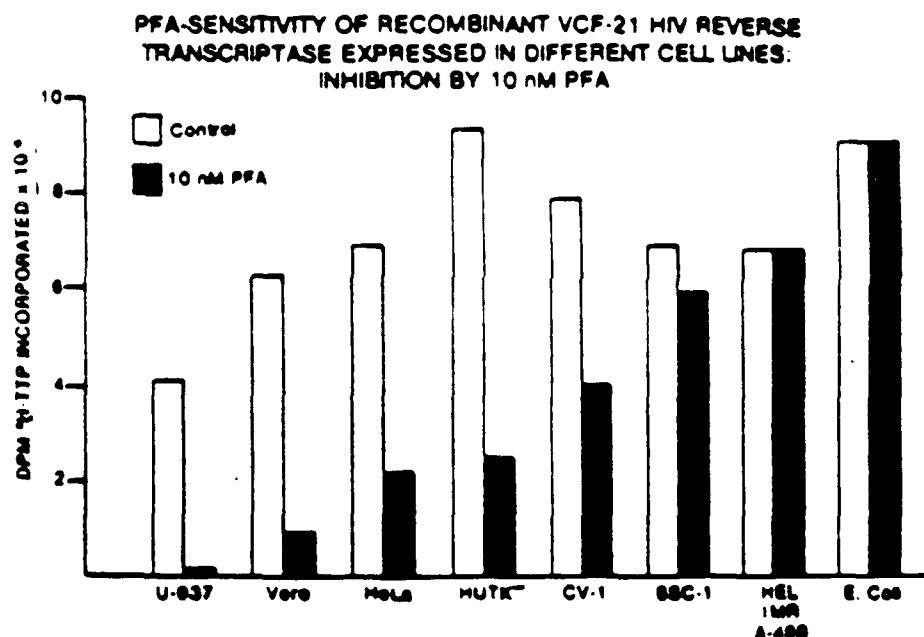
Figure: Inhibition of HTV-reverse transcriptase by synthetic AZT-triphosphate

Key O control ● .05 μ M □ 25 μ M Δ .25 μ M △ 2.5 μ M ■ Reagent blank

Preliminary experiments have been conducted to characterize further the nature of the inhibition by 3' uridine spiroxirane triphosphate. The inhibition is further confirmed as irreversible suicide type by the fact that addition of excess template does not reverse it, thus distinguishing it from that of a chain terminating inhibitor where new template would be expected to overcome the inhibition.

Expression of HIV-Reverse Transcriptase In Different Cell Lines.

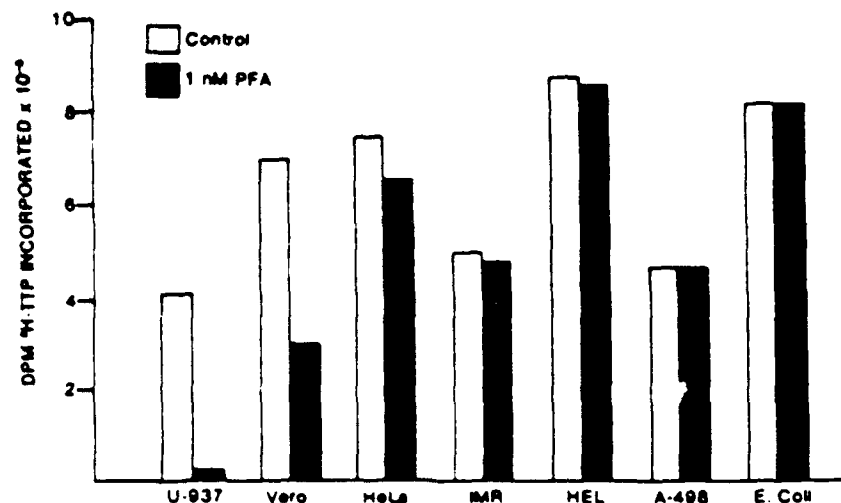
In order to determine if the recombinant HIV-reverse transcriptase was expressed in different forms depending upon the cell type, the vaccinia VCF-21 construct was grown in a number of different cell lines of both human, monkey and rodent origin. The expressed reverse transcriptase was tested for inhibition by Foscarnet at two different levels (1 and 10 nanomolar) and compared to the E. Coli recombinant HIV-RT (Kindly donated by Dr. Steven Hughes Fort Detrick M.D.) and the wild type HIV-RT. Both the wild type and E. Coli HIV-RT's were resistant to PFA showing essentially no inhibition at the 10nM level. Previous studies have shown that both enzymes have I_{50} 's for PFA in the 200-400 nM range. The recombinant HIV-RT expressed in eukaryotic cells however showed a range of phenotypes as indicated in figures 3 and 4 below. Both U-937 and Vero cells expressed enzyme sensitive to 1 nanomolar PFA, whereas human embryo lung and A-498 cells expressed RT-enzyme having wild-type sensitivity. Hela, HuTK- and CV-1 cells as observed previously expressed enzyme having intermediate PFA sensitivity



Sensitivity of Recombinant HIV-Reverse Transcriptases to 10 Nanomolar PFA.

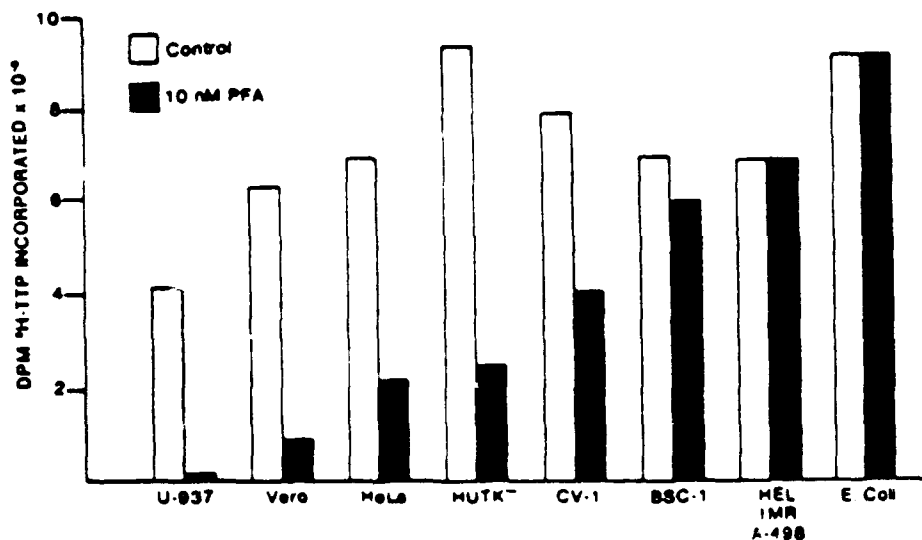
The VCF-21 vaccinia construct was grown in the indicated cell lines and the activity of the expressed reverse transcriptase was measured in the presence (dark blocks) or absence (open blocks) of 10 nanomolar PFA.

expressed RT-enzyme having wild-type sensitivity. Hela, HuTK- and CV-1 cells as observed previously expressed enzyme having intermediate PFA sensitivity.



Sensitivity of Recombinant HIV-Reverse Transcriptases to 1 Nanomolar PFA.

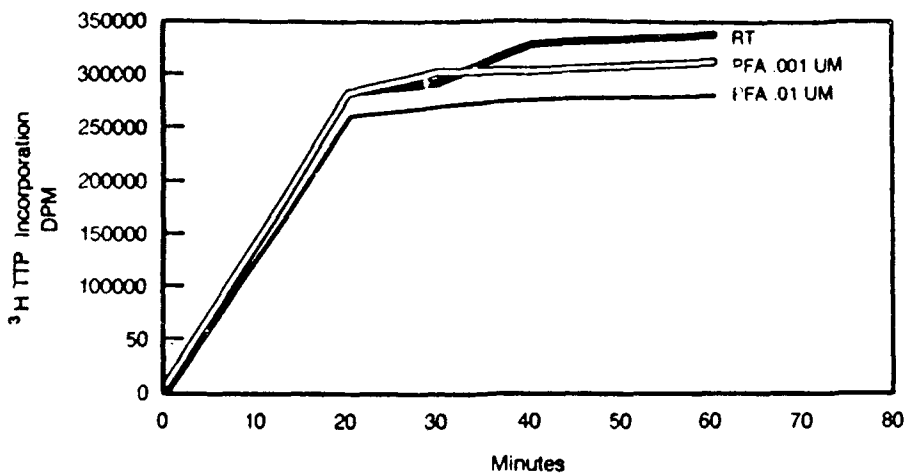
The VCF-21 vaccinia construct was grown in the indicated cell lines and the activity of the expressed reverse transcriptase was measured in the absence (open blocks) or presence of 1 nanomolar PFA. The height of the open blocks is a measure of the yield of recombinant enzyme activity in the different cell lines. Note the extreme sensitivity of the HIV-RT to PFA when expressed in the U-937 macrophage and vero monkey kidney cell lines.



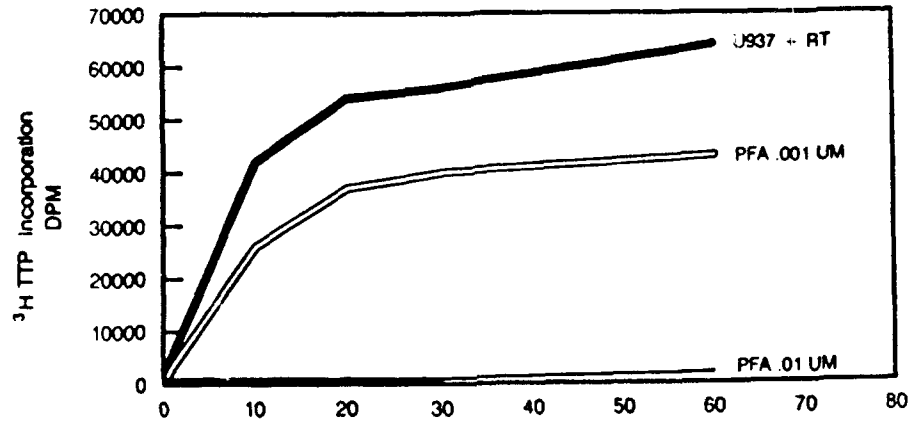
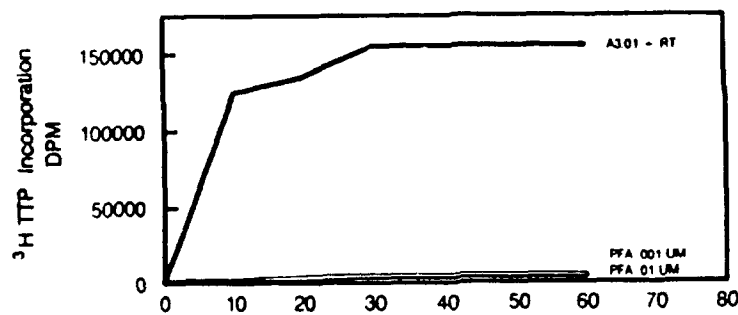
Sensitivity of Recombinant HIV-Reverse Transcriptases to 10 Nanomolar PFA.

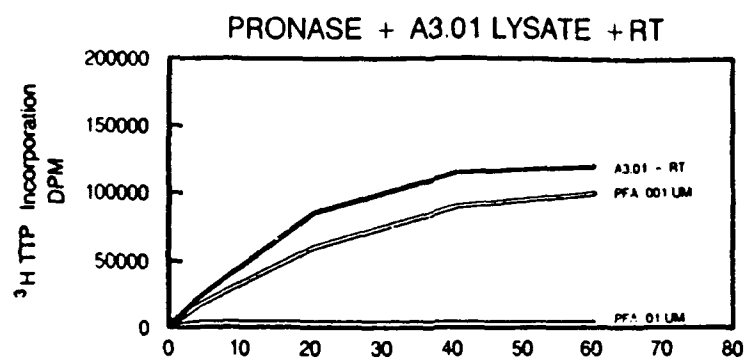
The VCF-21 vaccinia construct was grown in the indicated cell lines and the activity of the expressed reverse transcriptase was measured in the presence (dark blocks) or absence (open blocks) of 10 nanomolar PFA. Note that the enzyme produced in HEL, IMR and A498 cultures resembles the E. Coli recombinant and wild type HIV-RT in being relatively resistant to PFA. The enzyme expressed in HeLa, HVTK and CVI cells is of intermediate sensitivity and that expressed in U-937 and vero cells is maximally inhibited by PFA. Sensitivity of the vero enzyme remained following purification. Cross sensitizing experiments on mixtures of PFA sensitive and resistant preparations and measurements of RNase H activity of all constructs are in progress.

RECOMBINANT HIV RT (E. COLI) + FOSCARNET

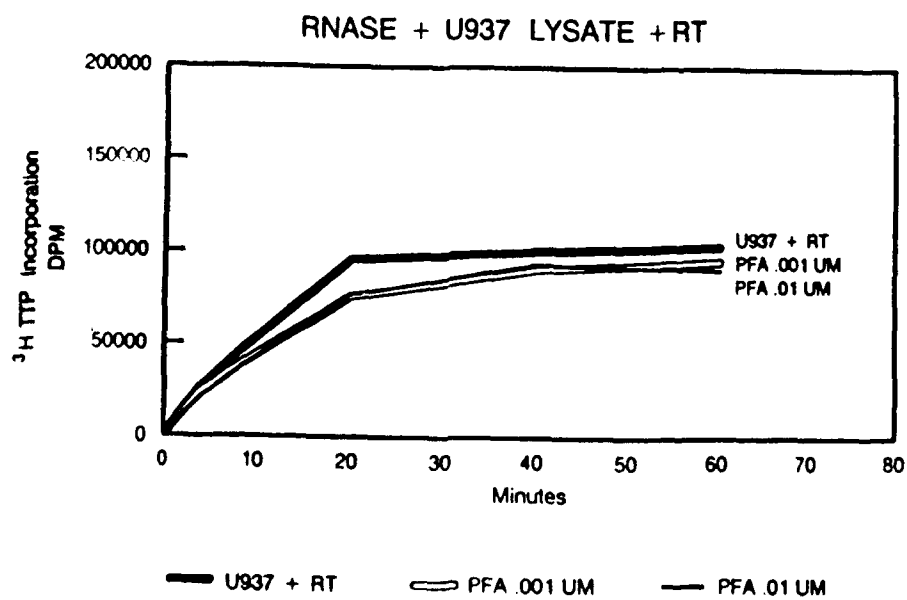
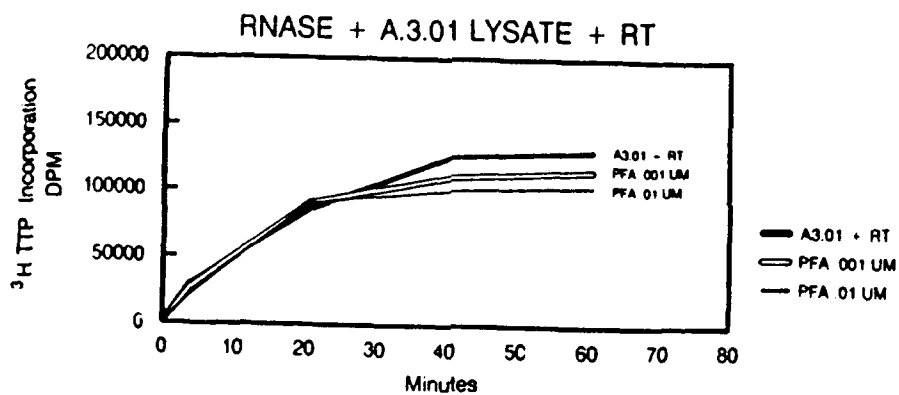


SENSITIVITY TO FOSCARNET INHIBITION OF HIV
REVERSE TRANSCRIPTASE (PURIFIED) IN THE
PRESENCE OF CELL LYSATES





**SENSITIVITY TO FOSCARNET INHIBITION OF HIV
REVERSE TRANSCRIPTASE (PURIFIED) IN THE
PRESENCE OF CELL LYSATES**



Appendix I

PUBLICATIONS

PROJECT: DAMD 17-87-C-7171

TITLE: SUICIDE INHIBITOR OF REVERSE TRANSCRIPTASE IN THERAPY OF AIDS AND OTHER RETROVIRUSES.

PRINCIPAL INVESTIGATOR: Dr. J.M. Bailey, Ph.D., D.Sc.
Professor of Biochemistry

PRODUCTIVITY REPORT

Publications:

1. Enhanced sensitivity to Foscarnet of first-strand viral replication by recombinant HIV-reverse transcriptase. M.M. Lightfoote and J.M. Bailey. FASEB J. 4:1318 (1990).
2. Differential sensitivity of wild-type and recombinant HIV-Reverse transcriptase to inhibition by Foscarnet. M.M. Lightfoote and J.M. Bailey. Proc Vth Int. AIDS Conf. Montreal. 5:515 (1989).
3. M.M. Lightfoote and J.M. Bailey. Somatic Cell Modulation of HIV-Reverse Transcriptase Expression. Antiviral Chem. and Chemotherapy. (1989) in preparation.
4. Nucleotide and template selectivity for inhibition of reverse transcriptase by PFA: Implications for retroviral therapy. J.M. Bailey and M.M. Lightfoote. Proc. IVth Int. AIDS Conf. Montreal. 4:3223 (1988).
5. Differential sensitivity of wild-type and recombinant HIV-reverse transcriptase to inhibition by foscarnet. M.M. Lightfoote and J.M. Bailey. Proc. IVth Int. AIDS Conf. Montreal. (1989).
6. Antiviral activities of some sterol phosphonoformate diester. J.M. Bailey, K. Nelson, M. Lightfoote. J. Clinic. Exp. Ther. in preparation.
7. Nucleoside spirox.ranes: A new class of retroviral inhibitor. J.M. Bailey, K. Nelson, M. Lightfoote. J. Virol. in preparation.
8. Synthesis and antiviral activities of some sterol dicarboxylate esters of 3' Azido thymidine (AZT). J.M. Bailey, R.M. Mook, M. Lightfoote. J. Clin. Exp. Ther. in preparation.
9. Synthesis of mono and di-substituted cholesterol phosphonoformates by the Arbuzov reaction. J.M. Bailey and Keith Nelson. Tetrahedron Letters. in preparation.
10. Drug sensitizing RNA interactions with HIV reverse transcriptase. J.M. Bailey and M. Lightfoote. Biochem. Soc. Trans. (1991) in press.

Appendix II
COMPOUNDS SYNTHESIZED:

Compounds synthesized and prepared for shipment to USAMRIID for antiviral testing.

1. 2',0²-Anhydouridine
2. 2',0²-Anhydrocytidine hydrochloride
3. 3',5'-Di-0-benzoyl-2'-0²-anhydouridine
4. 5'-0- β -Butyldimethylsilyl-3'-0-benzoyl-2',0'-anhydouridine
5. 2',3'-Anhydro-5'-0-trityluridine
6. 3'-Deoxy-2'-thymidinene
7. N³-Benzyl-2',5'-di-0-trityluridine
8. 5'-0- β -Butyldimethylsilylanhydrouridine
9. N⁴-Benzoyleytidine
10. 2',3'-Di-0-mesyl-5'-0-trityluridine
11. 5'-0- β -Butyldimethylsilyl-2',3'-isopropylideneuridine
12. 2',3'-Isopropylideneuridine
13. 2',3'-0-Sulfinyluridine
14. 2',3'-Benzylideneuridine
15. N⁴-Benzoyl-2',3'-0-Sulfinylcytidine
16. 2',3'-0-Sulfinylcytidine
17. 3',5'-Di-0-trityl-2'-deoxy-2'-oxouridine
18. 3',5'-Di-0- β -butyldimethylsilyl-2'-deoxy-2'-oxouridine
19. 2',5'-Di-0- β -butyldimethylsilyl-3'-deoxy-3'-oxouridine
20. Diethyl (cholesteryloxycarbonyl) phosphonate
21. Disodium (cholesteryloxycarbonyl) phosphonate
22. Di-[1-(3-carboethoxypropyl)] cholesteryloxycarbonyl
23. Di-(2,3-isopropylideneglycetyl) cholesteryloxycarbonyl phosphonate
24. Di-[1-(3-methylbutyl)] cholesteryloxycarbonyl phosphonate
Di-[1-(lithium 3-carboxypropyl)] cholesteryloxycarbonyl phosphonate
25. Sodium ethyl (cholesteryloxycarbonyl) phosphonate

26. Sodium 1-(3-carboxypropyl) 1-(30 carboethoxypropyl) [cholesteryloxycarbonyl] phosphonate
27. Adenosine 2',3'-Riboepoxide
28. Thymidine 5'-(1,3,2-dioxaphosphorin-2-oxide)
29. Thymidinene 5'-(1,3,2-dioxaphosphorin-2-oxide)
30. Thymidinene
31. 2-Ethoxy-5-chloro-6-methyl-1,3,2-dioxaphosphorin-5-ene-2-oxide
32. 2-Ethoxy-5-chloro-1,2-oxaphosphol-4-ene-2-oxide
33. 2,4-dichloro-5-methyl-1,3,2-dioxaphosphole-2-oxide
34. 2-methoxy-4,5-dimethyl-1,3,2-dioxaphole-2-oxide
35. Thymidine 3',5'-oxetane

Appendix III

TEST DATA ON ANTI HIV DRUGS

USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-II-55	AVS NO AVS-006466
		DATE RECD 12-28-89	AMT RECEIVED [mg]	MOL WT (au) 224.218
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME 2',3'-DIDEOXYTHYMIDINENE				

COMPOUND NAME

2', 3' -DIDEOXYTHYMIDINENE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

PLATE 1HK
DRUG 6466

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6466
TAI: 35.67 SI: 9.78

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.128	0.131	0.131	0.130	0.130	0.131	0.037	0.034	0.035	0.035	0.034	0.034
B	1.400	1.627	0.356	0.363	0.376	1.639					cc/vc	
C	1.362	1.625	0.418	0.386	0.384	1.622					1.651	
D	1.367	1.657	0.627	0.570	0.580	1.712					1.588	
E	1.430	0.369	1.821	1.729	1.850	1.785					1.666	
F	1.481	0.364	1.581	1.414	1.737	1.753					0.358	
G	0.160	0.368	0.127	0.130	0.142	0.156					0.343	
H	0.133	0.129	0.131	0.135	0.131	0.132					0.338	

tox-cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

CELLS

SHIPMENT NUMBER

STRN

REAGENT

VIRUS CONTROL

CELL CONTROL

DIFFERENTIAL

HIV3B

MT2 Satisfactory; Active; Retest

PROJECT #

6520-2

SPONSOR

USAMRIID

TEST DATE

04/04/90

DATE READ

04/12/90

	DRUG 6466	25%	50%	95%
TC (uG/mL)		48.80	66.40	97.90
IC (uG/mL)		3.53	4.99	9.33
ANTIVIRAL INDEX (AI)		13.84	13.29	10.49

DRUG 6466		ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% RED. IN VIRAL CPE	MEAN O.D.	% CELL VIABILITY			
low B	0.32	0.006	0%	1.387	92%			0.002
C	1	0.038	3%	1.361	90%			0.001
D	3.2	0.231	18%	1.404	93%			0.005
E	10	1.442	100%	1.476	98%			0.001
F	32	1.222	96%	1.488	99%			-0.001
high G	100	-0.227	0%	0.025	2%			0.003

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6466 vs. HIV3B (04/04/90)

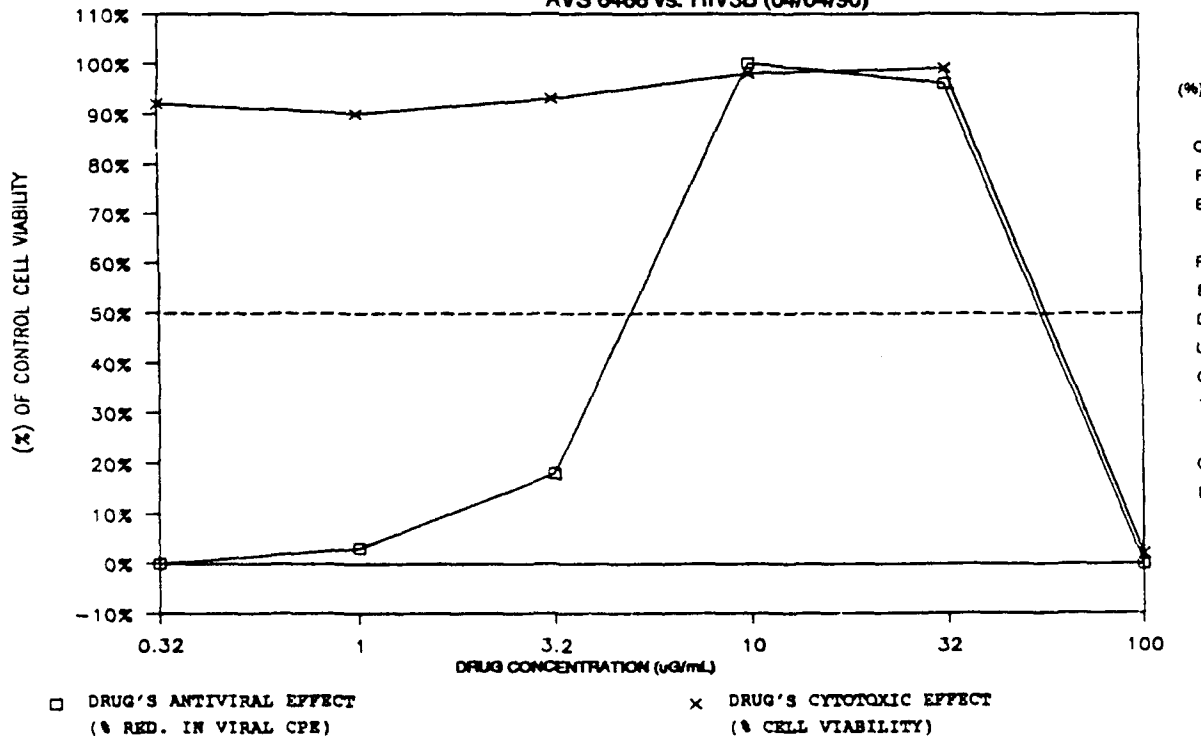


PLATE 1KA

IN VITRO ANTIVIRAL RESULTS

DRUG 6466

MTT ASSAY

DRUG: AVS 6466

TAI: >85.47 SI: >161.06

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background											
A	0.118	0.109	0.116	0.115	0.112	0.114	0.034	0.034	0.034	0.034	0.034	0.033
B	cc/vc 1.089						tox	drug 6466 experimental	cc/vc	tox		
C	1.051						1.190	1.183	1.176	1.067	1.233	1.133
D	0.917						1.137	1.101	1.193	1.236	1.212	1.245
E	0.478						1.061	1.207	1.204	1.234	1.205	1.323
F	0.469						1.197	1.195	1.334	1.246	0.512	1.477
G	0.500						1.629	1.371	1.380	1.360	0.576	1.578
H							0.302	0.174	0.174	0.154	0.566	0.248
	drug 6466 colorimetric background											
							0.144	0.137	0.135	0.126	0.121	0.132

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

HIVCRF

CELLS

CEM Satisfactory; Active; Retest

PROJECT # 6520-2

SHIPMENT NUMBER

64 Low MOI

SPONSOR USAMRIID

STRN

RP#

TEST DATE 04/26/90

REAGENT

0.114

DATE READ 05/03/90

VIRUS CONTROL

DRUG 6466

25%

50%

95%

CELL CONTROL

TC (uG/mL)

51.50

71.10

> 100.00

DIFFERENTIAL

IC (uG/mL)

ANTIVIRAL INDEX (AI)

> 161.06

> 222.13

> 312.50

DRUG 6466		ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% RED. IN CPE	MEAN O.D.	% CELL VIABILITY			
low B	0.32	0.606	100%	1.030	100%			0.018
C	1	0.652	100%	1.070	100%			0.007
D	3.2	0.685	100%	1.066	100%			0.012
E	10	0.720	100%	1.202	100%			0.021
F	32	0.830	100%	1.467	100%			0.023
high G	100	- .377	0%	0.131	13%			0.030

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6466 vs. HIVCRF (04/26/90)

The graph plots two metrics against drug concentration (0.32 to 100 uG/mL). The Y-axis represents (% of Control Cell Viability) from -10% to 110%. A horizontal dashed line at 50% indicates the threshold for cytotoxicity. The X-axis represents Drug Concentration (uG/mL) on a logarithmic scale. Data points are connected by lines:

- Antiviral Effect (Reduction in Viral CPE):** Represented by open squares. It remains near 100% until 32 uG/mL, then drops sharply to about 10% at 100 uG/mL.
- Cytotoxic Effect (Cell Viability):** Represented by crosses. It remains near 100% until 32 uG/mL, then drops sharply to about 10% at 100 uG/mL.

Drug Concentration (uG/mL)	Antiviral Effect (% Red. in Viral CPE)	Cytotoxic Effect (% Cell Viability)
0.32	100	100
1	100	100
3.2	100	100
10	100	100
32	100	100
100	~10	~10

DRUG'S ANTIVIRAL EFFECT
(% RED. IN VIRAL CPE)

DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

PRINTED 06/10/90

SOUTHERN RESEARCH INSTITUTE

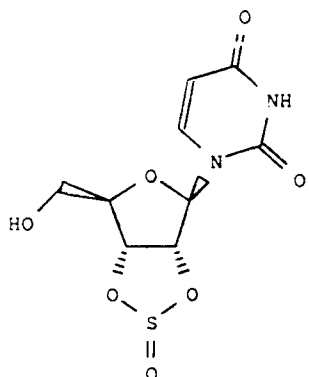
USAMRIID

Antiviral Drug Screening Program

03/26/90

STRUCTURE

CHIRAL



SUBMITTER 01141.01	CTR NO KN-V-99	AVS NO AVS-006442
DATE RECD 12-28-89	AMT RECEIVED [mg] 74.00	MOL WT (au) 290.253
HANDLING/STORAGE		
SOLUBILITY		
STABILITY		
ALT NAME 2',3'-O-SULFINYL URIDINE		

COMPOUND NAME

2',3'-O-SULFINYL URIDINE

SCREEN INSTRUCTION

PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD?>VSV

IN VIVO TOXICITY [mg/kg]

HOST VH RTE LD50 MTC LAB PR DATE

IN VITRO SCREEN [ug/ml]

VIR	VR	VR+	ID50	CELL	MTC	TI	TI+	LAB PRT DATE
JE	NOT ACT	VERO	183		0	SO MTT	90-03-01	
PT	100	VERO	170		2.39	SO MTT	90-03-01	
SF	NOT ACT	VERO	172		0	SO MTT	90-03-01	
VEE	NOT ACT	VERO	38.3		0	SO MTT	90-03-02	
YF	NOT ACT	VERO	171		0	SO MTT	90-03-01	

IN VIVO SCREEN [Dose = mg/kg]

VIR HST VR VR+ DOSE MTC VEH RTE D TOX SP L PR DATE

PLATE U9A
DRUG 6442

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6442
TAI: 15.38 SI: 1.70

	1	2	3	4	5	6	7	8	9	10	11	12
A	reagent background						plastic background					
A	0.042	0.041	0.040	0.042	0.039	0.042	0.001	0.001	0.001	0.001	0.001	0.001
B	0.898	0.951	0.392	0.299	0.337	0.956					0.746	
C	0.938	1.033	0.389	0.295	0.345	0.912					0.972	
D	1.011	1.139	0.419	0.423	0.434	1.020					0.800	
E	1.034	0.329	0.559	0.504	0.542	1.102					0.334	
F	1.115	0.389	0.663	0.626	0.656	1.181					0.342	
G	0.243	0.353	0.227	0.227	0.219	0.229					0.396	
H	0.049	0.038	0.038	0.037	0.038	0.039						

low-cell toxicity

no-cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

PT

CELLS

VERO

Satisfactory; Active; Retest

PROJECT # 5975-1

SHIPMENT NUMBER 63

SPONSOR USAMRIID

STRN

ADAMES

TEST DATE 03/01/90

REAGENT

0.041

DATE READ 03/09/90

VIRUS CONTROL

0.316

DRUG 6442

25%

50%

95%

CELL CONTROL

0.899

TC (uG/mL)

170.00

239.00

> 320.00

DIFFERENTIAL

0.583

IC (uG/mL)

22.20

100.00

ANTIVIRAL INDEX (AI)

7.65

2.39

DRUG 6442		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	1	-0.013	100%	0.888	99%	-0.002
C	3.2	-0.011	100%	0.887	99%	-0.003
D	10	0.072	88%	0.979	100%	-0.004
E	32	0.181	69%	1.030	100%	-0.003
F	100	0.294	50%	1.110	100%	-0.003
high G *	320	-0.141	100%	0.187	21%	0.008

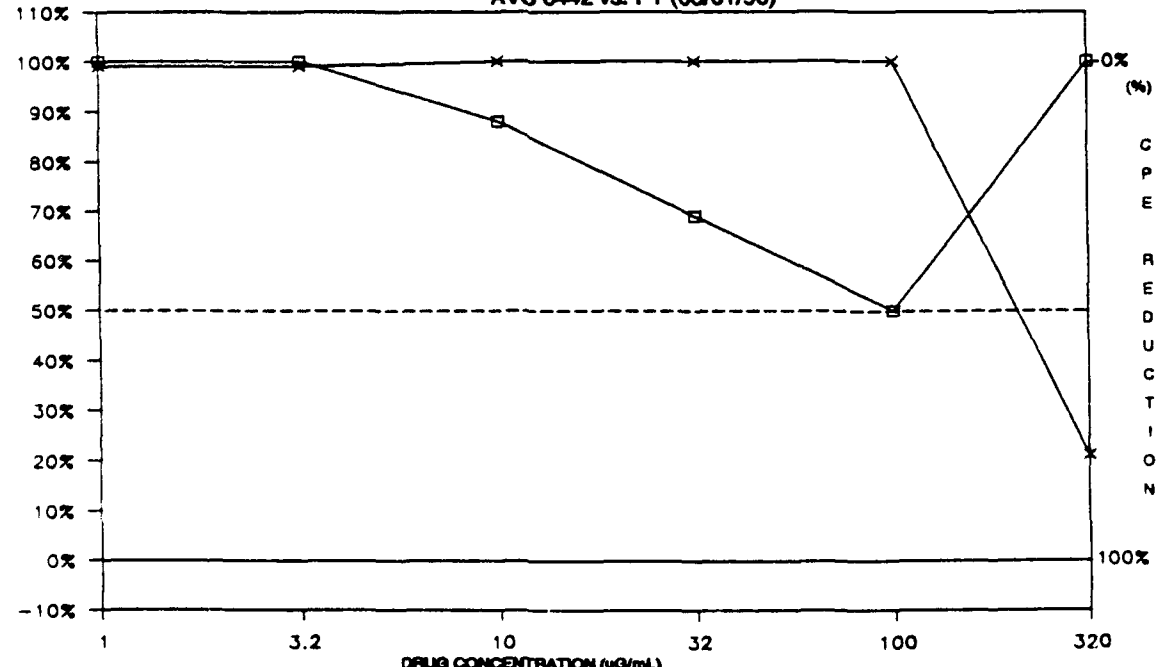
* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6442 vs. PT (03/01/90)

(%) OF CELL OR VIRUS CONTROL



□ DRUG'S ANTIVIRAL EFFECT
(% VIRAL CPE)

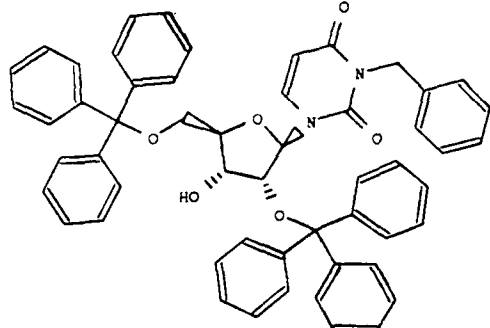
x DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

USAMRIID

Antiviral Drug Screening Program

03/26/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-V-109	AVS NO AVS-006443
		DATE RECD 12-28-89	AMT RECEIVED [mg] 86.00	MOL WT (au) 818.979
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME N3-BENZYL-2',5'-DI-O-TRITYLURIDINE				



COMPOUND NAME N3-BENZYL-2',5'-DI-O-TRITYLURIDINE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

IN VITRO SCREEN [ug/ml]	IN VIVO SCREEN [Dose = mg/kg]
VIR VR VR+ ID50 CELL MTC TI TI+ LAB PRT DATE	VIR HST VR VR+ DOSE MTC VEH RTE D TOX SP L PR DATE
JE NOT ACT VERO 24.7 0 SO MTT 90-03-01	
PT 77.1 VERO 210 > 4.15 SO MTT 90-03-01	
SF NOT ACT VERO > 320 0 SO MTT 90-03-01	
VEE NOT ACT VERO > 320 0 SO MTT 90-03-02	
YF NOT ACT VERO > 320 0 SO MTT 90-03-01	

PLATE U9A
DRUG 6443

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6443
TAI: >10.57 SI: 2.72

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plate background					
A	0.042	0.041	0.040	0.042	0.039	0.042	0.001	0.001	0.001	0.001	0.001	0.001
B		cells					tox	drug 6443 experimental	cells	tox		
B		0.951					0.830	0.376	0.357	0.334	0.746	0.879
C		1.033					0.911	0.386	0.324	0.297	0.972	0.771
D		1.139					1.015	0.406	0.436	0.343	0.800	0.866
E		0.329					0.800	0.493	0.491	0.497	0.334	0.814
F		0.389					0.734	0.695	0.713	0.683	0.342	0.716
G		0.353					0.696	0.560	0.632	0.599	0.396	0.751
H							drug 6443 colorimetric background					
							0.059	0.044	0.040	0.039	0.040	0.040

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

PT

CELLS

VERO Satisfactory; Active; Retest

PROJECT #

5975-1

SHIPMENT NUMBER

63

SPONSOR

USAMRIID

STRN

ADAMES

TEST DATE

03/01/90

REAGENT

0.041

DATE READ

03/09/90

VIRUS CONTROL

0.316

	DRUG 6443	25%	50%	95%
TC (uG/mL)		210.00	> 320.00	> 320.00
IC (uG/mL)		34.20	77.10	-----
ANTIVIRAL INDEX (AI)		6.15	> 4.15	-----

CELL CONTROL

0.899

DIFFERENTIAL

0.583

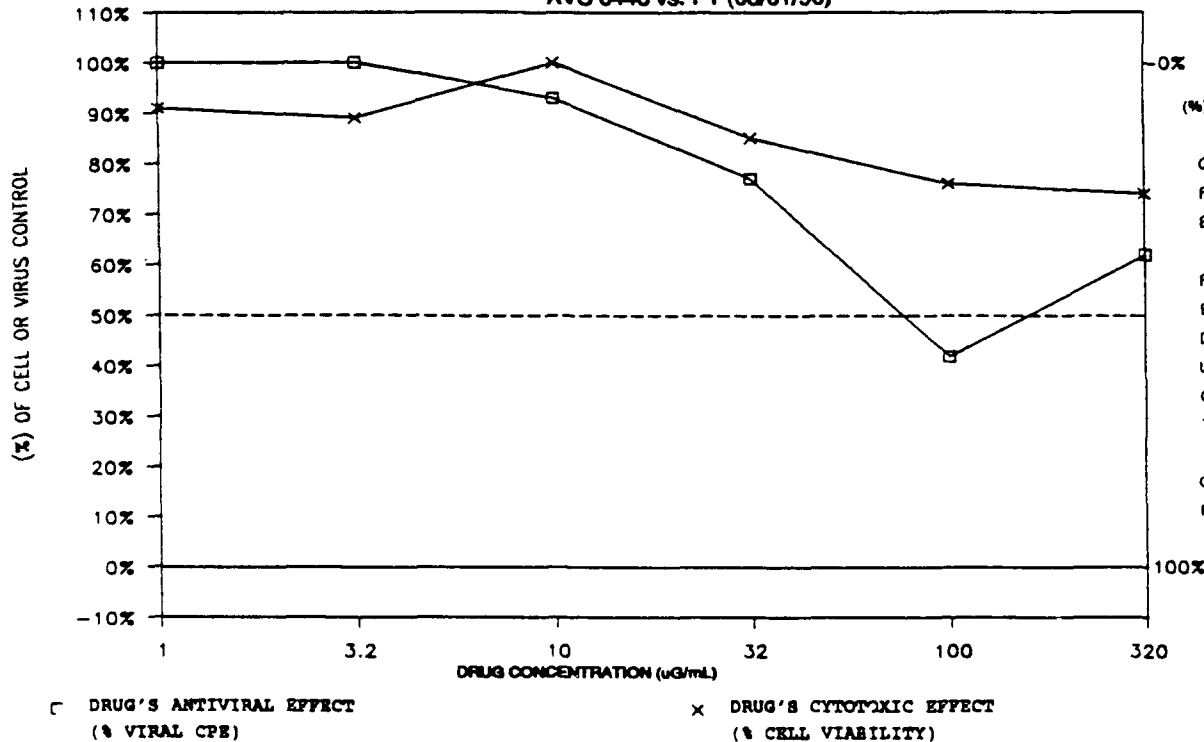
DRUG 6443		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	1	- .001	100%	0.815	91%	- .001
C	3.2	- .021	100%	0.801	89%	- .001
D	10	0.040	93%	0.902	100%	- .002
E	32	0.137	77%	0.767	85%	- .001
F	100	0.337	42%	0.681	76%	0.003
high G	320	0.222	62%	0.665	74%	0.018

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6443 vs. PT (03/01/90)

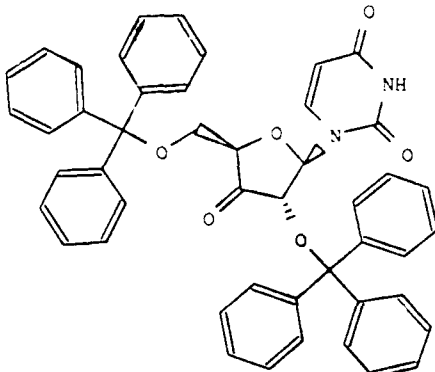


USAMRIID

Antiviral Drug Screening Program

03/26/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-VII-83	AVS NO AVS-006444
		DATE RECD 12-28-89	AMT RECEIVED [mg] 79.00	MOL WT (a.u.) 726.837
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME 3'-DEOXY-2',5'-DI-O-TRITYL-3'-OXOURIDINE				



COMPOUND NAME 3'-DEOXY-2',5'-DI-O-TRITYL-3'-OXOURIDINE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

IN VITRO SCREEN [ug/ml]	IN VIVO SCREEN [Dose = mg/kg]
VIR VR VR+ IO50 CELL MTC TI TI+ LAB PRT DATE	VIR HST VR VR+ DOSE MTC VEH RTE D TOX SP L PR DATE
JE NOT ACT VERO 51 0 SO MTT 90-03-01 PT 79.2 VERO > 320 > 4.04 SO MTT 90-03-01 SF NOT ACT VERO 30 0 SO MTT 90-03-01 VEE NOT ACT VERO > 320 0 SO MTT 90-03-02 YF NOT ACT VERO > 320 0 SO MTT 90-03-01	

PLATE U9B

IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6444
TAI: >24.65 SI: >4.04

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.043	0.042	0.045	0.041	0.040	0.042	0.001	0.001	0.001	0.001	0.001	0.001
B	1.033	0.918	0.365	0.330	0.321	0.864					0.837	
C	0.959	0.946	0.304	0.333	0.381	0.999					0.734	
D	0.936	1.054	0.433	0.451	0.414	0.883					0.631	
E	0.811	0.295	0.382	0.411	0.490	0.810					0.339	
F	0.753	0.355	0.577	0.706	0.655	0.748					0.369	
G	0.982	0.382	0.952	0.924	0.879	0.981					0.358	
H	drug 6444 colorimetric background											
	0.041	0.043	0.042	0.039	0.039	0.039						

toxic-cell toxicity

control control

virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS	PT	PROJECT #	5975-1
CELLS	VERO	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	03/01/90
STRN	ADAMES	DATE READ	03/09/90
REAGIFT	0.042	DRUG 64.4	25% 50% 95%
VIRUS CONTROL	0.308	TC (ug/mL)	> 320.00 > 320.00 > 320.00
CELL CONTROL	0.811	IC (ug/mL)	41.50 79.20 278.00
DIFFERENTIAL	0.504	ANTIVIRAL INDEX (AI)	> 7.72 > 4.04 > 1.15

DRUG 6444		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (ug/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	1	-.008	100%	0.909	100%	-.003
C	3.2	-.007	100%	0.940	100%	-.003
D	10	0.086	83%	0.870	100%	-.003
E	32	0.078	85%	0.768	35%	0.000
F	100	0.295	41%	0.707	87%	0.001
high G	320	0.570	0%	0.940	100%	-.001

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6444 vs. PT (03/01/90)

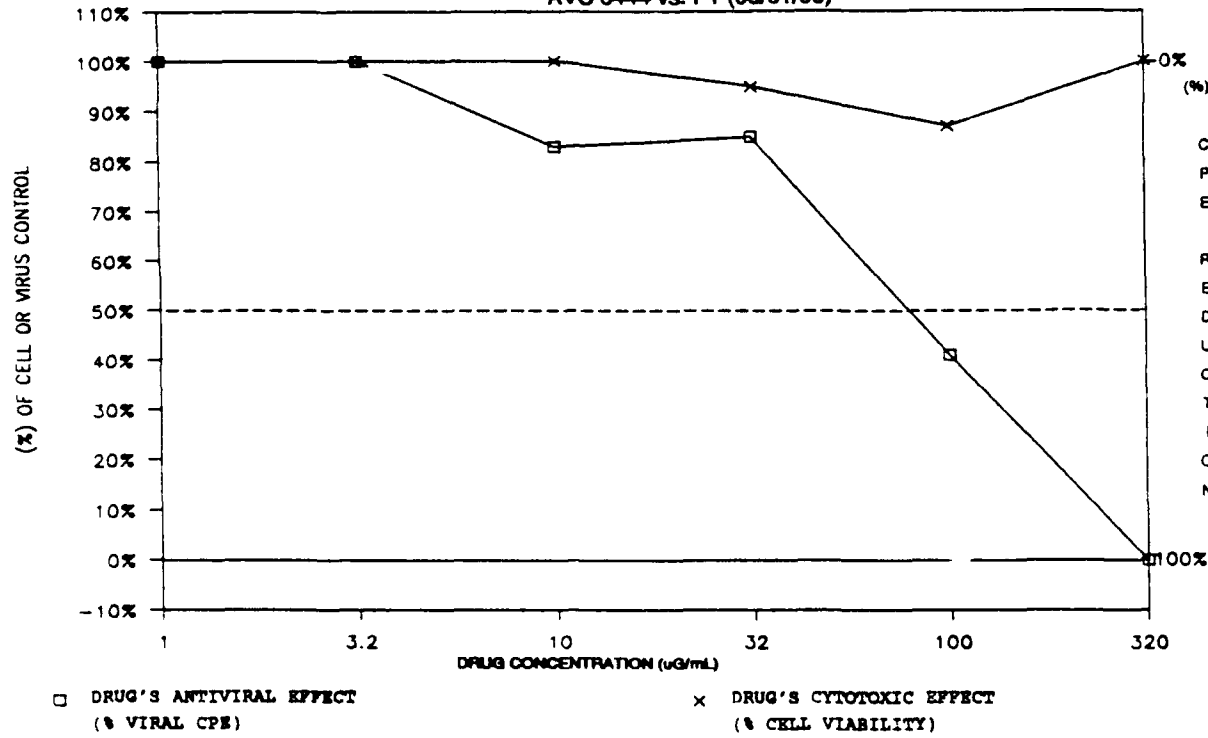


PLATE UAR
DRUG 6444

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6444
TAI: >7.30 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.042	0.040	0.039	0.038	0.039	0.039	0.001	0.001	0.001	0.001	0.001	0.001
tox	cc/vro		drug 6444 experimental			tox					cc/vro	
B	0.957	0.847	0.187	0.272	0.243	0.896					0.926	
C	0.960	1.048	0.275	0.297	0.280	1.005					0.782	
D	0.807	0.922	0.352	0.319	0.346	0.906					0.882	
E	0.817	0.213	0.421	0.441	0.493	0.717					0.218	
F	0.728	0.253	0.278	0.294	0.369	0.731					0.176	
G	0.972	0.297	0.071	0.062	0.069	0.798					0.190	
drug 6444 colorimetric background												
H	0.042	0.042	0.043	0.041	0.041	0.039						

tox=cell toxicity cc=cell control vc=virus control BOLD = highest drug conc values shown are optical densities

VIRUS	YF	PROJECT #	5975-1
CELLS	VERO	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	03/01/90
STRN	ASIBI	DATE READ	03/09/90
REAGENT	0.040	DRUG 6444	25% 50% 95%
VIRUS CONTROL	0.185	TC (ug/mL)	> 320.00 > 320.00 > 320.00
CELL CONTROL	0.862	IC (ug/mL)	17.90 ----- -----
DIFFERENTIAL	0.677	ANTIVIRAL INDEX (AI)	> 17.89 ----- -----

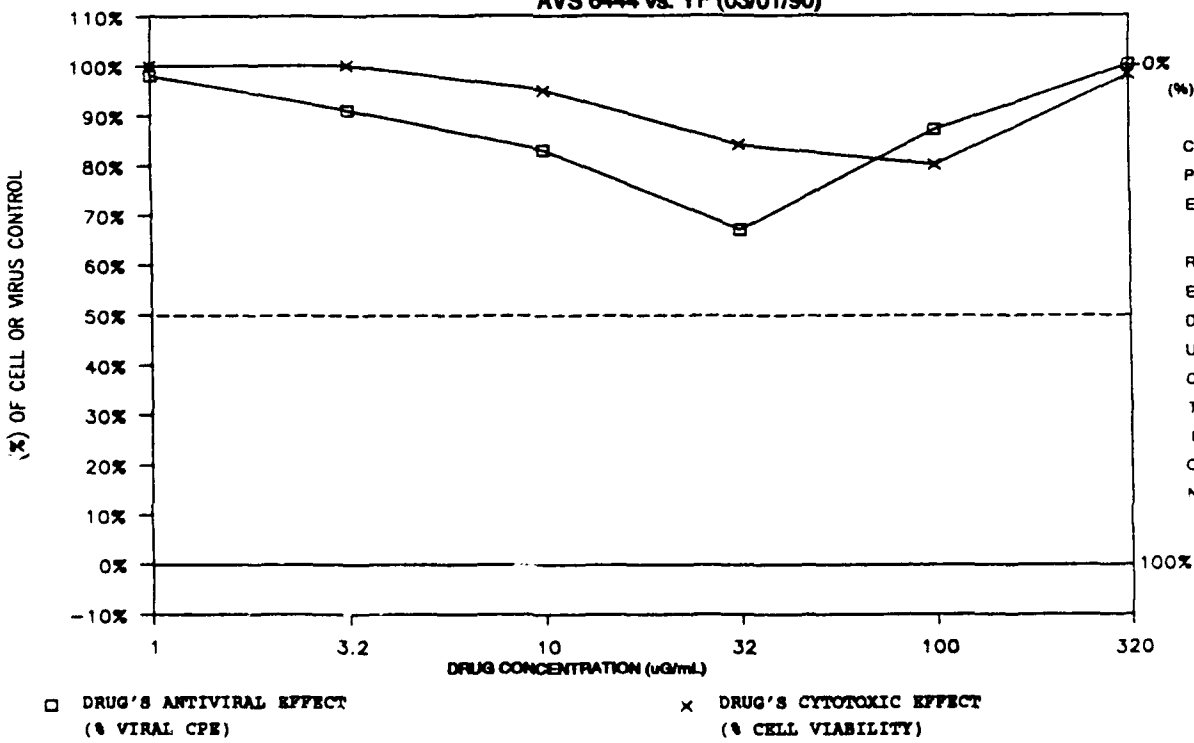
ROW ON PLATE	CONC. (ug/mL)	ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	MEAN O.D.	% VIABILITY	
low B	1	0.011	98%	0.888	100%	0.888	100%	-.001
C	3.2	0.058	91%	0.941	100%	0.941	100%	0.002
D	10	0.113	83%	0.815	95%	0.815	95%	0.002
E	32	0.224	67%	0.725	84%	0.725	84%	0.003
F	100	0.086	87%	0.687	80%	0.687	80%	0.003
high G *	320	-.160	100%	0.843	98%	0.843	98%	0.003

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6444 vs. YF (03/01/90)



USAMRIID

Antiviral Drug Screening Program

03/26/90

STRUCTURE	CHIRAL	SUBMITTER	CTR NO	AVS NO
		01141.01	KN-VII-21	AVS-006445
		DATE RECD 12-28-89	AMT RECEIVED [mg] 74.00	MOL WT (au) 726.837
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME 2'-DEOXY-3',5'-DI-O-TRITYL-2'-OXOURIDINE				

COMPOUND NAME

2'-DEOXY-3',5'-DI-O-TRITYL-2'-OXOURIDINE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

IN VITRO SCREEN [ug/ml]	IN VIVO SCREEN [Dose = mg/kg]
VIR VR VR+ ID50 CELL MTC TI TI+ LAB PRT DATE JE NOT ACT VERO 23.2 0 SO MTT 90-03-01 PT 22.6 VERO 49 2.92 SO MTT 90-03-01 SF NOT ACT VERO 43.3 0 SO MTT 90-03-01 VEE NOT ACT VERO 32 0 SO MTT 90-03-02 YF 18.9 VERO 48 3.45 SO MTT 90-03-01	VIR HST VR VR+ DOSE MTC VEH RTE D TOX SP L PR DATE

PLATE U9B
DRUG 6445

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6445
TAI: >17.14 SI: 2.17

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.043	0.042	0.045	0.041	0.040	0.042	0.001	0.001	0.001	0.001	0.001	0.001
B		co/va					tox	drug 6445 experimental	co/va	tox		
C		0.918					0.833	0.402	0.373	0.379	0.837	0.871
D		0.946					1.074	0.286	0.430	0.407	0.734	0.870
E		1.054					0.830	0.587	0.567	0.534	0.631	0.816
F		0.295					0.882	0.656	0.606	0.590	0.339	0.890
G		0.355					0.033	0.036	0.038	0.035	0.369	0.033
H		0.382					0.035	0.036	0.036	0.036	0.358	0.037
							drug 6445 colorimetric background					
							0.038	0.044	0.043	0.040	0.039	0.039

tox=cell toxicity

co=cell control

vo=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

PT

PROJECT #

5975-1

CELLS

VERO

Satisfactory; Active; Retest

SPONSOR

USAMRIID

SHIPMENT NUMBER

63

TEST DATE

03/01/90

STRN

ADAMES

DATE READ

03/09/90

REAGENT

0.042

DRUG 6445

25%

50%

95%

VIRUS CONTROL

0.308

TC (uG/mL)

49.00

66.00

96.60

CELL CONTROL

0.811

IC (uG/mL)

5.74

22.60

DIFFERENTIAL

0.504

ANTIVIRAL INDEX (AI)

8.53

2.92

DRUG 6445		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	1	0.038	92%	0.813	100%	- .003
C	3.2	0.028	94%	0.933	100%	- .003
D	10	0.215	57%	0.783	97%	- .002
E	32	0.267	47%	0.843	100%	0.001
F	100	.315	100%	.311	0%	0.002
high G	320	.310	100%	.002	0%	- .004

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6445 vs. PT (03/01/90)

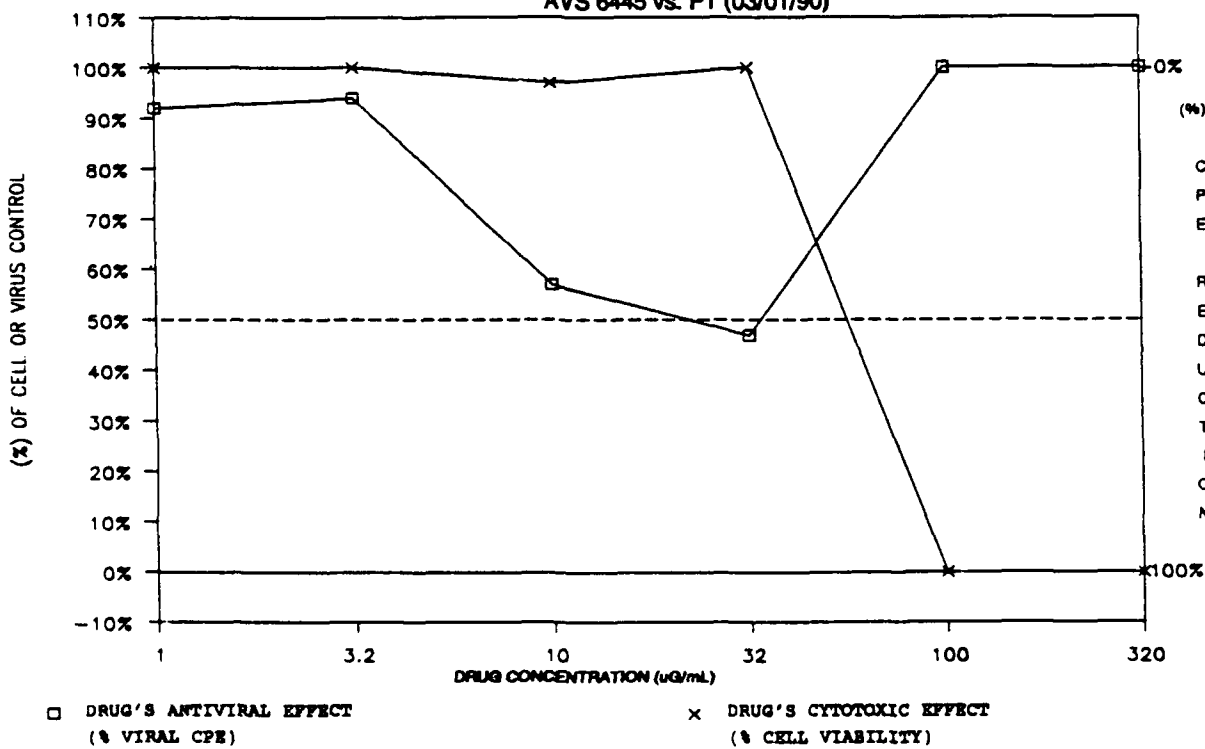


PLATE UAR
DRUG 6445

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6445
TAI: >19.99 SI: 2.53

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.042	0.040	0.039	0.038	0.039	0.039	0.001	0.001	0.001	0.001	0.001	0.001
B		0.048					tox	drug 6445 experimental	cc/vc	tox		
C		0.048					0.912	0.290	0.356	0.317	0.926	0.943
D		0.922					0.867	0.433	0.430	0.426	0.782	0.925
E		0.213					0.767	0.426	0.441	0.474	0.882	0.834
F		0.253					0.857	0.621	0.700	0.670	0.218	0.916
G		0.297					0.034	0.033	0.034	0.034	0.176	0.042
H							0.035	0.036	0.035	0.035	0.190	0.038
							drug 6445 colorimetric background					
							0.038	0.041	0.044	0.039	0.040	0.042

tox=cell toxicity cc=cell control vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS	YF	PROJECT #	5975-1
CELLS	VERO	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	03/01/90
STRN	ASIBI	DATE READ	03/09/90
REAGENT	0.040	DRUG 6445	258
VIRUS CONTROL	0.185	TC (ug/ml)	48.00
CELL CONTROL	0.862	IC (ug/ml)	2.22
DIFFERENTIAL	0.677	ANTIVIRAL INDEX (AI)	21.56
			508
			958

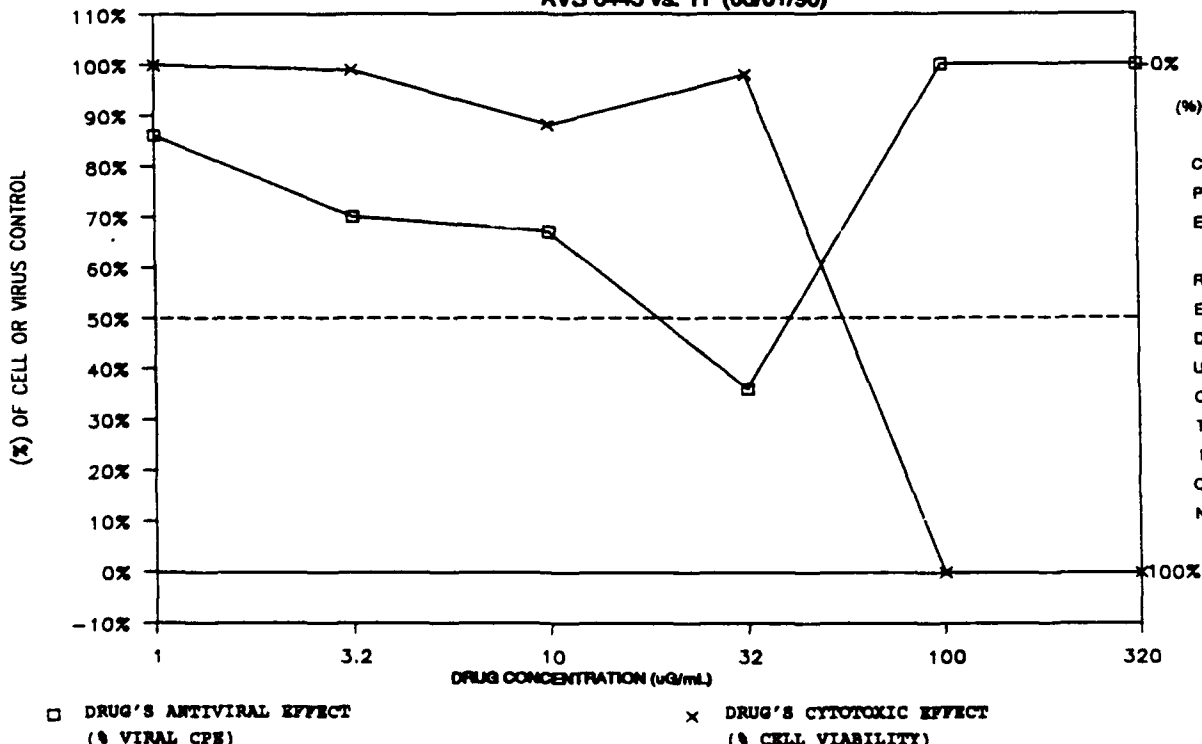
ROW ON PLATE	CONC. (ug/ml)	ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY			
low B	1	0.094	86%	0.885	100%			0.003
C	3.2	0.204	70%	0.856	99%			0.001
D	10	0.224	67%	0.762	88%			-.001
E	32	0.435	36%	0.843	98%			0.004
F	100	-.193	100%	-.003	0%			0.002
high G *	320	-.187	100%	-.001	0%			-.002

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6445 vs. YF (03/01/90)



Appendix IV
TEST DATA AGAINST OTHER VIRUSES

USAMRIID

Antiviral Drug Screening Program

09/21/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-II-95	AVS NO AVS-006467	
		DATE RECD 12-28-89	AMT RECEIVED [mg] 72.60	MOL WT (au) 261.667	
HANDLING/STORAGE					
SOLUBILITY					
STABILITY					
ALT NAME 2',O2-ANHYDROCYTIDINE HYDROCHLORIDE					

COMPOUND NAME

2',_O2-ANHYDROCYTIDINE HYDROCHLORIDE

PLATE 1HK
DRUG 6467

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6467
TAI: 0.00 SI: _____

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.128	0.131	0.131	0.130	0.130	0.131	0.037	0.034	0.035	0.035	0.034	0.034
B	1.627						tox	drug 6467 experimental			1.651	0.184
C	1.625						0.172	0.157	0.155	0.147	1.588	0.138
D	1.657						0.132	0.125	0.130	0.130	1.666	0.139
E	0.369						0.131	0.128	0.130	0.133	0.358	0.143
F	0.364						0.138	0.135	0.131	0.133	0.343	0.130
G	0.368						0.137	0.132	0.133	0.131	0.338	0.140
H							0.135	0.134	0.132	0.132	0.130	0.132
							drug 6467 colorimetric background					
							0.125	0.129	0.131	0.129	0.130	0.132

tox=cell toxicity cc=cell control vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS	HIV3B	PROJECT #	6520-2
CELLS	MT2	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	04/04/90
STRW	2.5	DATE READ	04/12/90
REAGENT	0.130	DRUG: 6467	25% 50% 95%
VIRUS CONTROL	0.227	TC (uG/mL)	< 0.32 < 0.32 < 0.32
CELL CONTROL	1.506	IC (uG/mL)	-----
DIFFERENTIAL	1.279	ANTIVIRAL INDEX (AI)	-----

ROW ON PLATE	CONC. (uG/mL)	ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
		MEAN O.D.	% RED. IN CPE	MEAN O.D.	% CELL VIABILITY			
low B	0.32	-.206	0%	0.046	3%			0.002
C	1	-.228	0%	0.005	0%			0.000
D	3.2	-.225	0%	0.006	0%			-.001
E	10	-.225	0%	0.009	1%			0.001
F	32	-.224	0%	0.004	0%			-.001
high G	100	-.219	0%	0.012	1%			-.005

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6467 vs. HIV3B (04/04/90)

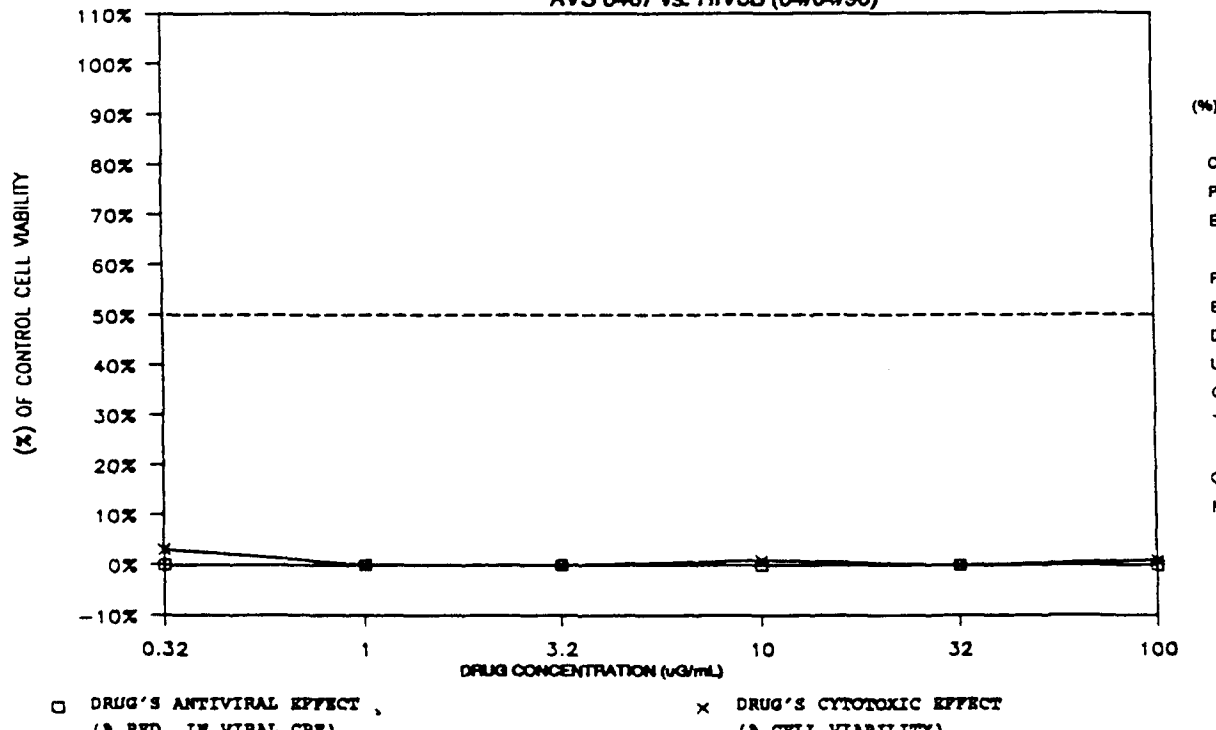


PLATE 1JP
DRUG 6467

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6467
TAI: >0.42 SI: —

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background											
A	0.117	0.112	0.113	0.115	0.114	0.116	0.034	0.035	0.035	0.035	0.035	0.034
	tox	cc/vc	drug 6467 experimental			tox					cc/vc	
B	1.266	1.279	0.277	0.333	0.325	1.350					1.389	
C	1.184	1.290	0.307	0.305	0.307	1.295					1.346	
D	1.306	1.275	0.320	0.311	0.315	1.371					1.311	
E	1.114	0.289	0.257	0.245	0.268	1.244					0.280	
F	0.628	0.300	0.216	0.199	0.240	0.715					0.207	
G	0.124	0.301	0.119	0.124	0.119	0.124					0.294	
	drug 6467 colorimetric background											
H	0.116	0.113	0.113	0.113	0.112	0.115						

tox=cell toxicity cc=cell control vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS	HIV3B	PROJECT #	6520-2
CELLS	MT2	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	04/24/90
STRN	2.5	DATE READ	05/02/90
REAGENT	0.115	DRUG: 6467	25% 50% 95%
VIRUS CONTROL	0.164	TC: (uG/mL)	0.040000 0.093600 0.922000
CELL CONTROL	1.201	IC: (uG/mL)	— — —
DIFFERENTIAL	1.037	ANTIVIRAL INDEX (AI)	— — —

ROW ON PLATE	CONC. (nG/mL)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% RED. IN VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	0.01	0.032	3%	1.193	99%	0.001
C	0.1	0.031	3%	1.128	94%	-.003
D	1	0.039	4%	1.226	100%	-.002
E	10	-.020	0%	1.067	89%	-.002
F	100	-.058	0%	0.559	47%	-.002
high G	1000	-.160	0%	0.007	1%	0.002

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6467 vs. HIV3B (04/24/90)

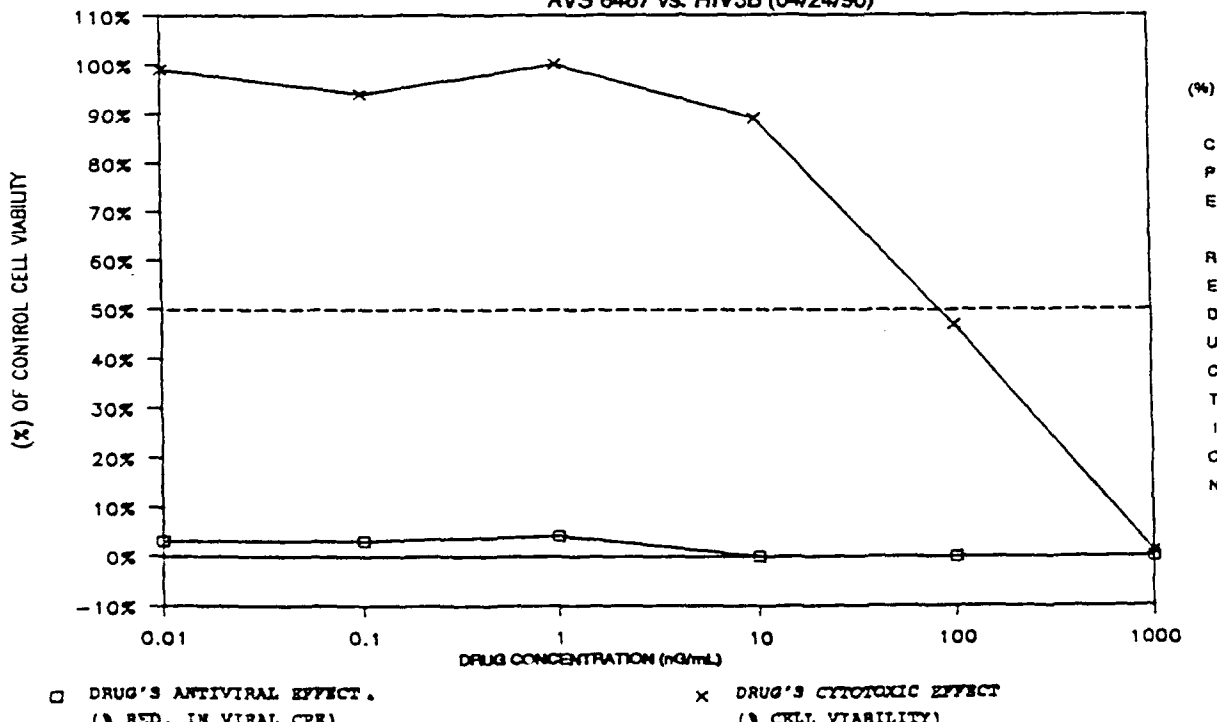


PLATE 0U4
DRUG 6467

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6467
TAI: 23.24 SI: 4.65

	1	2	3	4	5	6	7	8	9	10	11	12	
A	0.105	0.097	reagent background 0.123	0.114	0.109	0.119	0.000	0.000	0.000	0.000	0.000	0.000	
B		active 1.314					tox 1.488	0.153	0.166	0.134	1.321	1.330	
C		1.339					1.510	0.179	0.148	0.205	1.470	1.424	
D		1.312					1.553	0.258	0.206	0.325	1.410	1.427	
E		0.182					1.415	1.153	1.398	1.302	0.160	1.541	
F		0.180					1.058	0.847	0.883	0.733	0.173	0.825	
G		0.174					0.450	0.426	0.446	0.427	0.142	0.400	
H								drug 6467 colorimetric background 0.107	0.110	0.105	0.108	0.114	0.117

tox=cell toxicity cc=cell control vv=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS VV
CELLS VERO
SHIPMENT NUMBER 63
STRN LEDCA
REAGENT 0.111
VIRUS CONTROL 0.057
CELL CONTROL 1.250
DIFFERENTIAL 1.193

Satisfactory; Active; Retest
RETEST AT 3.2 UG/ML
PROJECT # 5975-4
SPONSOR USAMRIID
TEST DATE 04/19/90
DATE READ 04/25/90

DRUG 6467	25%	50%	95%
TC (uG/mL)	0.82	1.86	3.20
IC (uG/mL)	0.13	0.18	---
ANTIVIRAL INDEX (AI)	6.52	10.53	---

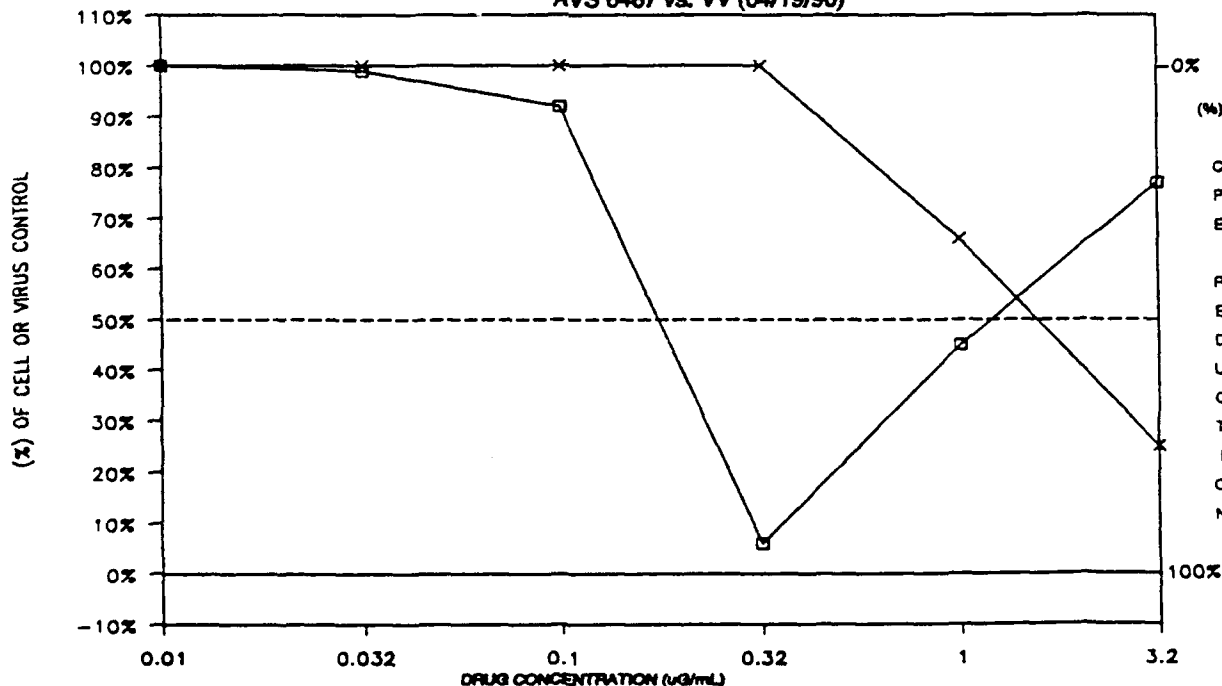
DRUG 6467		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN 0.D.	% VIRAL CPE	MEAN 0.D.	% CELL VIABILITY	
low B	0.01	-.023	100%	1.292	100%	0.006
C	0.032	0.006	99%	1.353	100%	0.003
D	0.1	0.098	92%	1.382	100%	-.003
E	0.32	1.122	6%	1.373	100%	-.006
F	1	0.654	45%	0.831	66%	-.001
high G *	3.2	0.269	77%	0.318	25%	-.004

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6467 vs. VV (04/19/90)



DRUG'S ANTIVIRAL EFFECT.
(% VIRAL CPE)

DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

PLATE OVV
DRUG 6467IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6467
TAI: 17.08 SI: 3.66

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.108	0.113	0.110	0.114	0.111	0.121	0.000	0.000	0.000	0.000	0.000	0.000
B		cc/vc					tox	drug 6467 experimental	cc/vc		tox	
C		1.830					1.845	0.203	0.233	0.193	1.728	1.710
D		1.920					1.733	0.174	0.147	0.218	1.917	1.735
E		1.895					1.913	0.357	0.263	0.292	1.915	1.868
F		0.199					1.843	1.232	1.174	1.173	0.136	1.739
G		0.124					1.495	1.117	1.250	1.167	0.166	1.229
H		0.186					0.497	0.416	0.431	0.448	0.110	0.556
							drug 6467 colorimetric background					
							0.104	0.106	0.109	0.104	0.116	0.109

tox=cell toxicity cc=cell control vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

VV

CELLS

VERO

Satisfactory

PROJECT #

5975-4

SHIPMENT NUMBER

63

CONFIRMS ORIGINAL ACTIVITY

SPONSOR

USAMRIID

STRM

LEDCA

TEST DATE

05/10/90

REAGENT

0.113

DRUG 6467

25%

50%

95%

VIRUS CONTROL

0.041

TC (uG/mL)

0.92

2.01

> 3.20

CELL CONTROL

1.755

IC (uG/mL)

0.14

0.35

DIFFERENTIAL

1.714

ANTIVIRAL INDEX (AI)

6.40

8.03

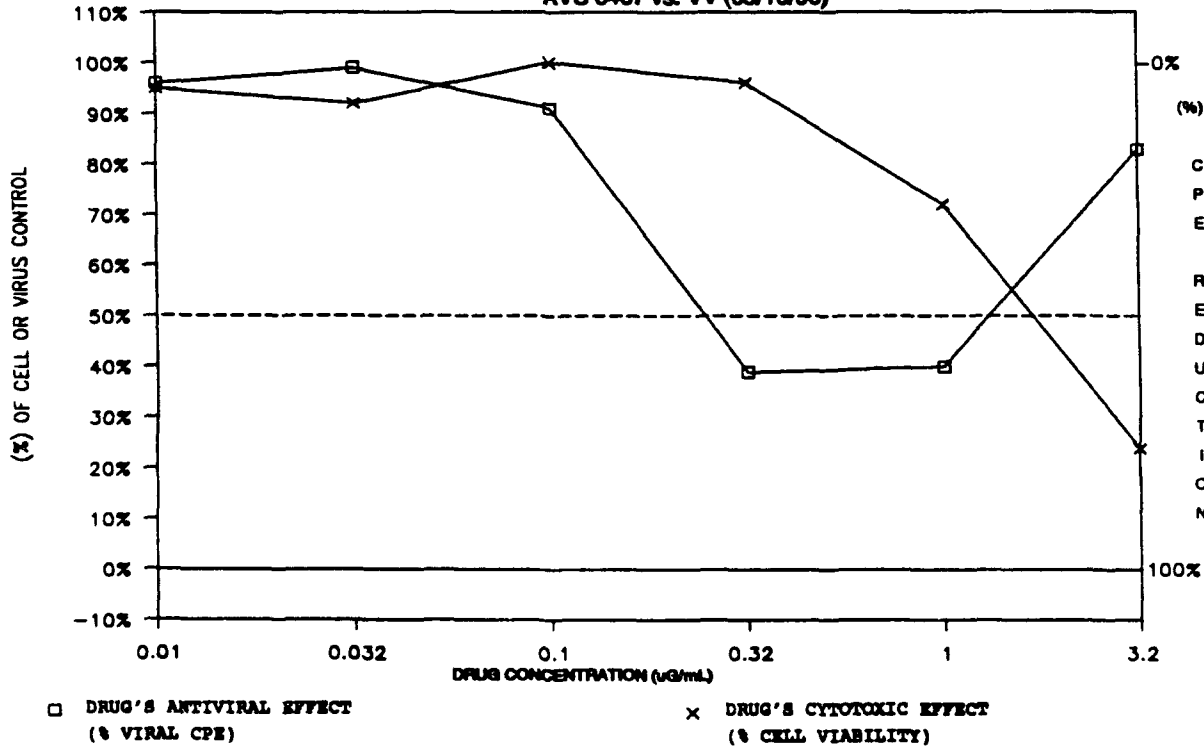
DRUG 6467		ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES				
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	COLORIMETRIC CONTROL			
low B	0.01	0.060	96%	1.669	95%				- .004
C	0.032	0.023	99%	1.618	92%				0.003
D	0.1	0.160	91%	1.787	100%				- .009
E	0.32	1.044	39%	1.682	96%				- .004
F	1	1.032	40%	1.256	72%				- .007
high G *	3.2	0.287	83%	0.423	24%				- .009

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6467 vs. VV (05/10/90)



USAMRIID

Antiviral Drug Screening Program

09/21/90

STRUCTURE	CHIRAL	SUBMITTER	CTR NO	AVS NO
		01141.01	KN-II-71	AVS-006462
		DATE RECD 12-28-89	AMT RECEIVED [mg] 72.40	MOL WT (au) 325.729
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME 2',3'-O-SULFINYL CYTIDINE HYDROCHLORIDE				

COMPOUND NAME

2',3'-O-SULFINYL CYTIDINE HYDROCHLORIDE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

PLATE 1HI
DRUG 6462

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6462
TAI: 0.00 SI: —

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.137	0.135	0.136	0.136	0.137	0.138	0.035	0.033	0.034	0.034	0.035	0.042
B	0.327	1.655	0.203	0.202	0.208	0.323					cave	
C	0.165	1.702	0.139	0.144	0.143	0.151					1.743	
D	0.139	1.663	0.130	0.133	0.130	0.137					1.651	
E	0.141	0.352	0.141	0.140	0.136	0.142					1.644	
F	0.138	0.358	0.132	0.132	0.131	0.133					0.346	
G	0.133	0.350	0.123	0.126	0.127	0.131					0.368	
H	0.127	0.130	0.132	0.131	0.134	0.135					0.337	

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

HIV3B

CELLS

MT2 Satisfactory; Toxic; Retest

SHIPMENT NUMBER

63

STRN

2.5

REAGENT

0.137

VIRUS CONTROL

0.215

CELL CONTROL

1.540

DIFFERENTIAL

1.325

PROJECT #

6520-2

SPONSOR

USAMRIID

TEST DATE

04/04/90

DATE READ

04/12/90

	DRUG 6462	25%	50%	95%
TC (ug/mL)	< 0.32	< 0.32	< 0.32	0.80
IC (ug/mL)	-----	-----	-----	-----
ANTIVIRAL INDEX (AI)	-----	-----	-----	-----

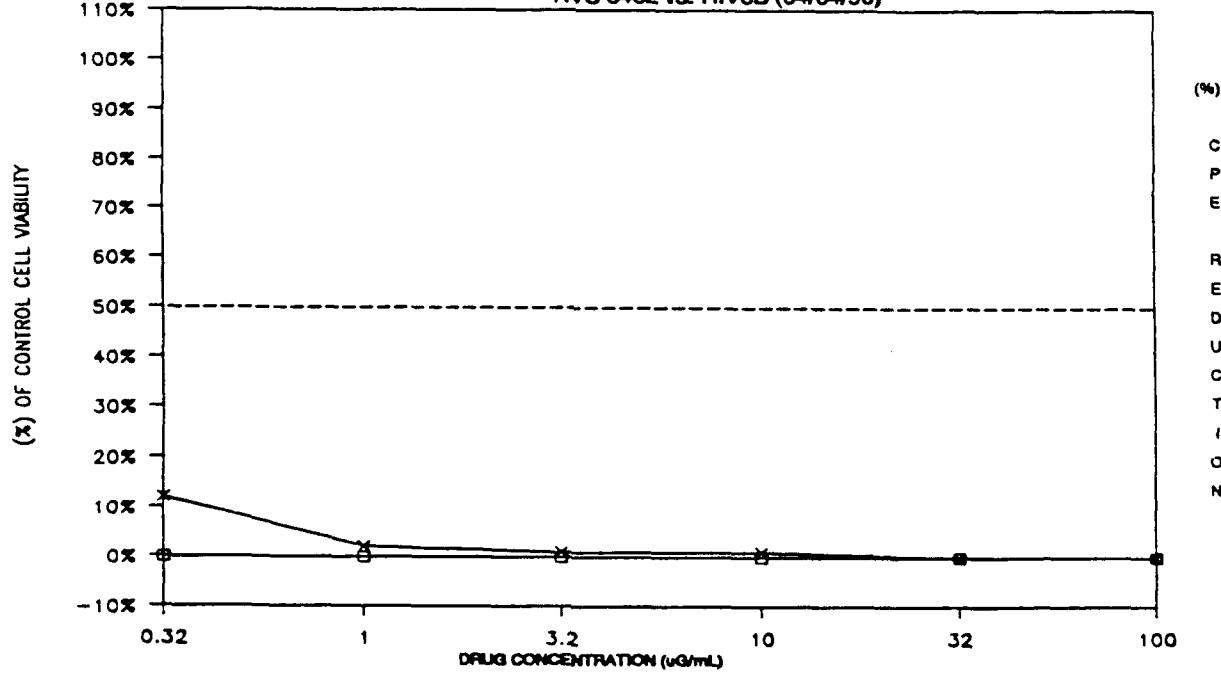
ROW ON PLATE	CONC. (uG/mL)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% RED. IN VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	0.32	-.146	0%	0.191	12%	-.002
C	1	-.207	0%	0.024	2%	-.003
D	3.2	-.215	0%	0.008	1%	-.006
E	10	-.208	0%	0.010	1%	-.005
F	32	-.213	0%	0.006	0%	-.007
high G	100	-.217	0%	0.005	0%	-.010

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6462 vs. HIV3B (04/04/90)



□ DRUG'S ANTIVIRAL EFFECT
(% RED. IN VIRAL CPE)

✖ DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

PLATE 1JO
DRUG 6462IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6462
TAI: 0.10 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.099	0.096	0.104	0.107	0.101	0.100	0.034	0.034	0.035	0.034	0.033	0.034
B		active 1.277					low	drug 6462 experimental	active	low		
C		1.183					1.202	0.325	0.302	0.296	1.184	1.192
D		1.477					1.204	0.317	0.328	0.336	1.249	1.164
E		0.309					1.277	0.318	0.322	0.327	1.201	1.297
F		0.288					1.193	0.320	0.302	0.316	0.271	1.098
G		0.266					0.906	0.218	0.264	0.275	0.303	0.842
H							0.125	0.121	0.118	0.122	0.292	0.119
							0.117	0.115	0.115	0.111	0.108	0.113

low=cell toxicity active control vs=virus control

BOLD = highest drug conc.

values shown are optical densities

VIRUS	HIV3B	PROJECT #	6520-2
CELLS	MT2	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	04/24/90
STRN	2.5	DATE READ	05/02/90
REAGENT	0.101	DRUG: 6462	25% 50% 55%
VIRUS CONTROL	0.187	IC: (uG/mL)	0.062500
CELL CONTROL	1.161	IC: (uG/mL)	0.308000
DIFFERENTIAL	0.974	ANTIVIRAL INDEX (AVI)	0.931000

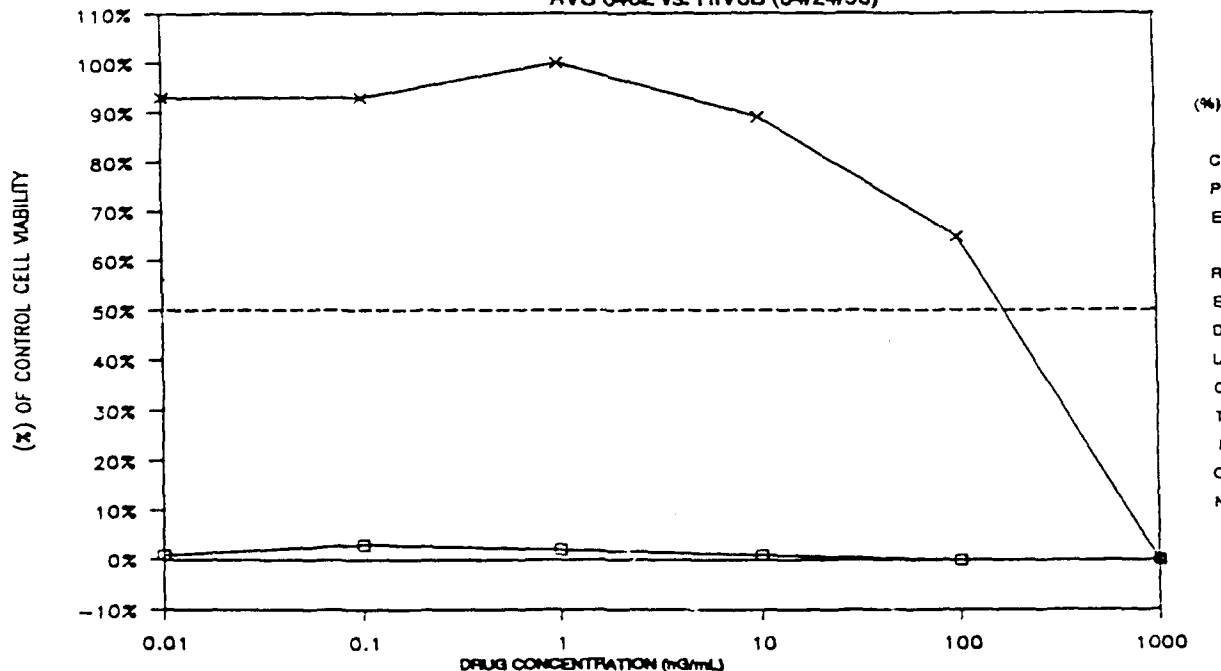
ROW ON PLATE	CONC. (nG/mL)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% RED. IN CPE	MEAN O.D.	% CELL VIABILITY	
low B	0.01	0.008	1%	1.084	93%	0.012
C	0.1	0.032	3%	1.076	93%	0.007
D	1	0.024	2%	1.176	100%	0.010
E	10	0.011	1%	1.030	89%	0.014
F	100	-0.050	0%	0.759	65%	0.014
high G	1000	-0.184	0%	0.005	0%	0.016

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6462 vs. HIV3B (04/24/90)



DRUG'S ANTIVIRAL EFFECT
(* RED. IN VIRAL CPE)

DRUG'S CYTOTOXIC EFFECT
(* CELL VIABILITY)

PLATE 0U2
DRUG 6462IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6462
TAI: 30.00 SI: 9.69

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.125	0.117	0.119	0.120	0.124	0.130	0.000	0.000	0.000	0.000	0.000	0.000
B	1.284	1.387	0.125	0.155	0.151	1.443					1.322	
C	1.197	1.494	0.453	0.332	0.473	1.315					1.435	
D	1.183	1.466	1.131	1.258	1.282	1.466					1.513	
E	1.311	0.176	1.234	1.216	1.347	1.456					0.278	
F	0.415	0.176	0.359	0.399	0.430	0.446					0.187	
G	0.280	0.197	0.263	0.256	0.255	0.294					0.173	
H	0.124	0.106	0.110	0.108	0.103	0.152						

tox=cell toxicity cc=cell control vo=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS
CELLS
SHIPMENT NUMBER
STRN
REAGENTVV
VERO Satisfactory
63 CONFIRMS ORIGINAL ACTIVITY
LEDCA
0.123PROJECT # 5975-4
SPONSOR USAMRIID
TEST DATE 04/19/90
DATE READ 04/25/90VIRUS CONTROL
CELL CONTROL
DIFFERENTIAL

	DRUG 6462	25%	50%	95%
VIRUS CONTROL	TC (uG/mL)	16.70	24.48	> 100.00
CELL CONTROL	IC (uG/mL)	1.10	1.72	-----
DIFFERENTIAL	ANTIVIRAL INDEX (AI)	15.27	14.12	-----

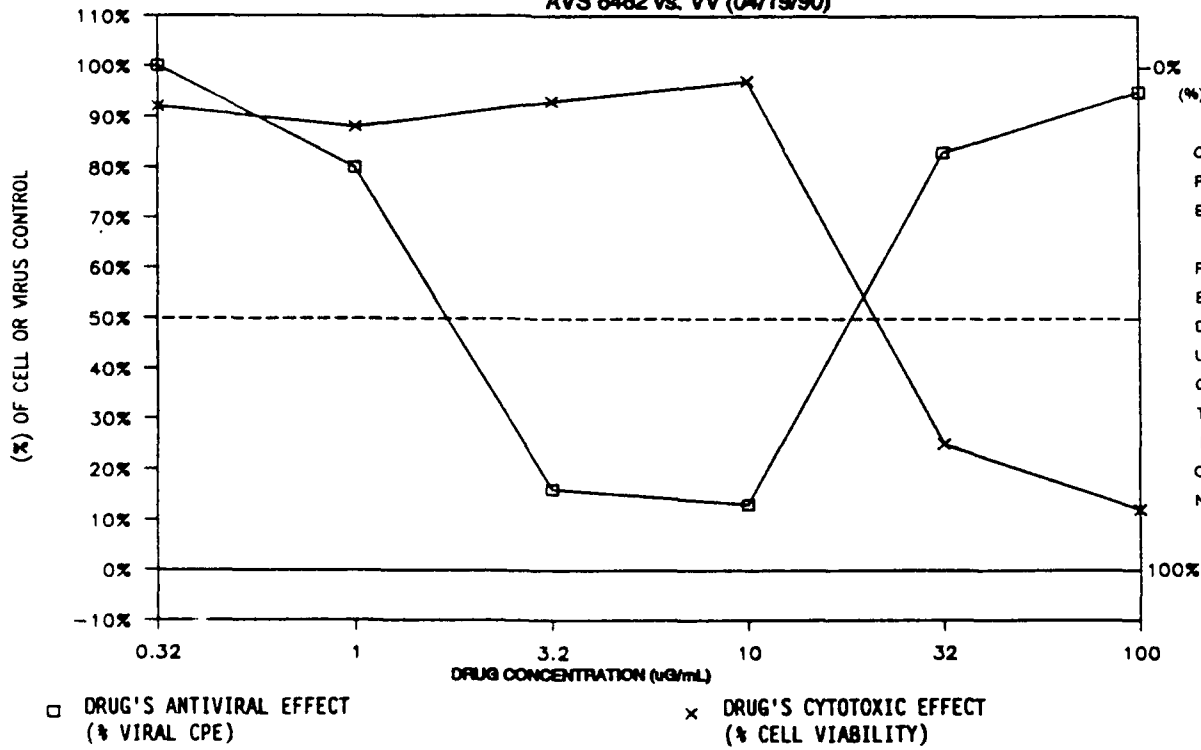
DRUG 6462		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL	
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	MEAN O.D.	% COLORIMETRIC CONTROL
low B	0.32	.084	100%	1.211	92%	0.029	
C	1	0.242	80%	1.154	88%	-0.020	
D	3.2	1.041	16%	1.217	93%	-0.015	
E	10	1.081	13%	1.274	97%	-0.013	
F	32	0.215	83%	0.325	25%	-0.017	
high G *	100	0.059	95%	0.164	12%	0.001	

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6462 vs. VV (04/19/90)



USAMRIID

Antiviral Drug Screening Program

29/21/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-VII-87	AVS NO AVS-006460
	DATE RECD 12-28-89	AMT RECEIVED [mg] 73.00	MOL WT (av) 740.864	
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME				
3'-DEOXY-2',5'-DI-O-TRITYL-3'-SPIROEPOXYURIDINE				

COMPOUND NAME

3'-DEOXY-2'-, 5'-DT-O-TRITYL-3' -SPIROEPOXYURIDINE

PLATE 1Q9
DRUG 6460IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6460
TAI: >30.24 SI: 2.79

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.302	0.273	0.281	0.414	0.402	0.296	0.079	0.081	0.295	0.252	0.130	0.071
B		1.940					1.999	0.798	0.734	0.680	2.096	2.160
C		1.881					1.993	0.697	0.802	0.720	1.992	1.693
D		1.776					1.987	0.947	0.994	0.879	1.856	1.943
E		0.557					2.067	1.159	0.949	0.911	0.423	2.060
F		0.423					2.018	0.930	1.338	1.309	0.365	1.921
G		0.478					1.223	0.719	0.614	0.637	0.500	0.881
H							0.354	0.270	0.282	0.268	0.273	0.313

tox=cell toxicity

co-cell control

vo=virus control

BOLD = highest drug conc.

values shown are optical densities

VIRUS	HIVCRF	PROJECT #	6520-2
CELLS	CEM	SPONSOR	USAIRRIID
SHIPMENT NUMBER	63	TEST DATE	06/12/90
STRN	RF2	DATE READ	06/19/90
REAGENT	0.328	IC (uG/ml)	254
VIRUS CONTROL	0.130	IC (uG/ml)	82.48
CELL CONTROL	1.596	IC (uG/ml)	1.18
DIFFERENTIAL	1.466	ANTIVIRAL INDEX (AI)	52.81
			954
			100.00

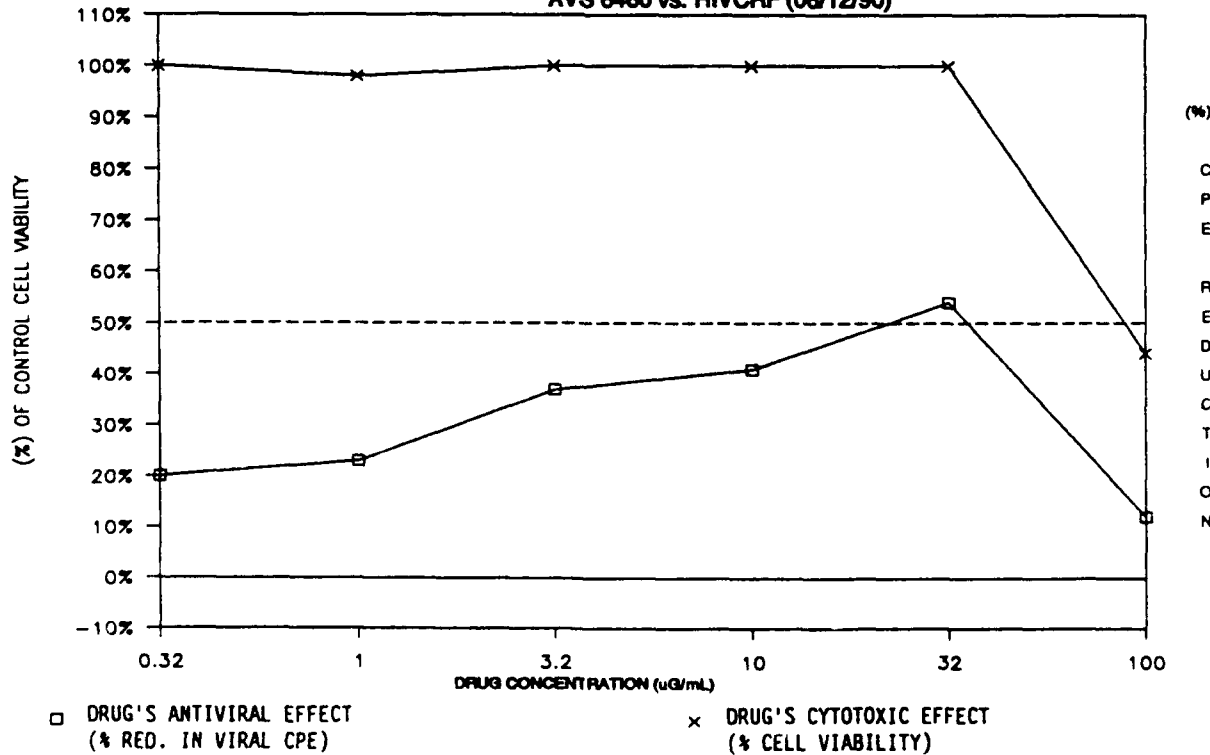
DRUG 6460		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/ml)	MEAN O.D.	% RED. IN CPE	MEAN O.D.	% CELL VIABILITY	
Tow B	0.32	0.295	20%	1.767	100%	- .015
C	1	0.337	23%	1.570	98%	- .055
D	3.2	0.542	37%	1.697	100%	- .060
E	10	0.595	41%	1.782	100%	- .046
F	32	0.793	54%	1.700	100%	- .058
high G *	100	0.173	12%	0.698	44%	0.026

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6460 vs. HIVCRF (06/12/90)



USAMRIID

Antiviral Drug Screening Program

39/21/30

STRUCTURE	SUBMITTER 01141.01	CTR NO MS-I-48	AVS NO AVS-006457
	DATE RECD 12-28-89	AMT RECEIVED [mg] 71.40	MOL WT (au) 538.621
HANDLING/STORAGE			
SOLUBILITY			
STABILITY			
ALT NAME DISODIUM(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE			

COMPOUND NAME

DISODIUM(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

PLATE 1Q8
DRUG 6457IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6457
TAI: >27.78 SI: >1.29

	1	2	3	4	5	6	7	8	9	10	11	12
			reagent background						plastic background			
A	0.433	0.373	0.552	0.655	0.447	0.451	0.103	0.100	0.191	0.442	0.164	0.087
^{tox}	^{above}	^{drug 6457 experimental}	^{tox}								^{above}	
B	1.739	1.747	0.707	0.601	0.807	1.972					1.841	
C	1.606	1.765	0.674	0.715	1.069	1.794					1.750	
D	1.871	1.913	0.559	0.799	1.037	1.806					2.012	
E	1.782	0.522	0.965	0.990	0.742	1.916					0.811	
F	1.783	0.541	0.894	0.662	1.126	1.899					0.583	
G	1.691	0.602	1.069	1.161	1.241	1.811					0.652	
H	0.345	0.405	0.387	0.396	0.393	0.382						
	tox-cell toxicity	co-cell control	vo-virus control				BOLD = highest drug conc					values shown are optical densities

VIRUS	HIVCRF	PROJECT #	6520-2
CELLS	CEM	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	06/12/90
STRN	RF2	DATE READ	06/19/90
REAGENT	0.485	DRUG 6457	25%
VIRUS CONTROL	0.133	IC (uG/ml)	> 100.00
CELL CONTROL	1.353	IC (uG/ml)	4.28
DIFFERENTIAL	1.220	ANTIVIRAL INDEX (AI)	> 21.37

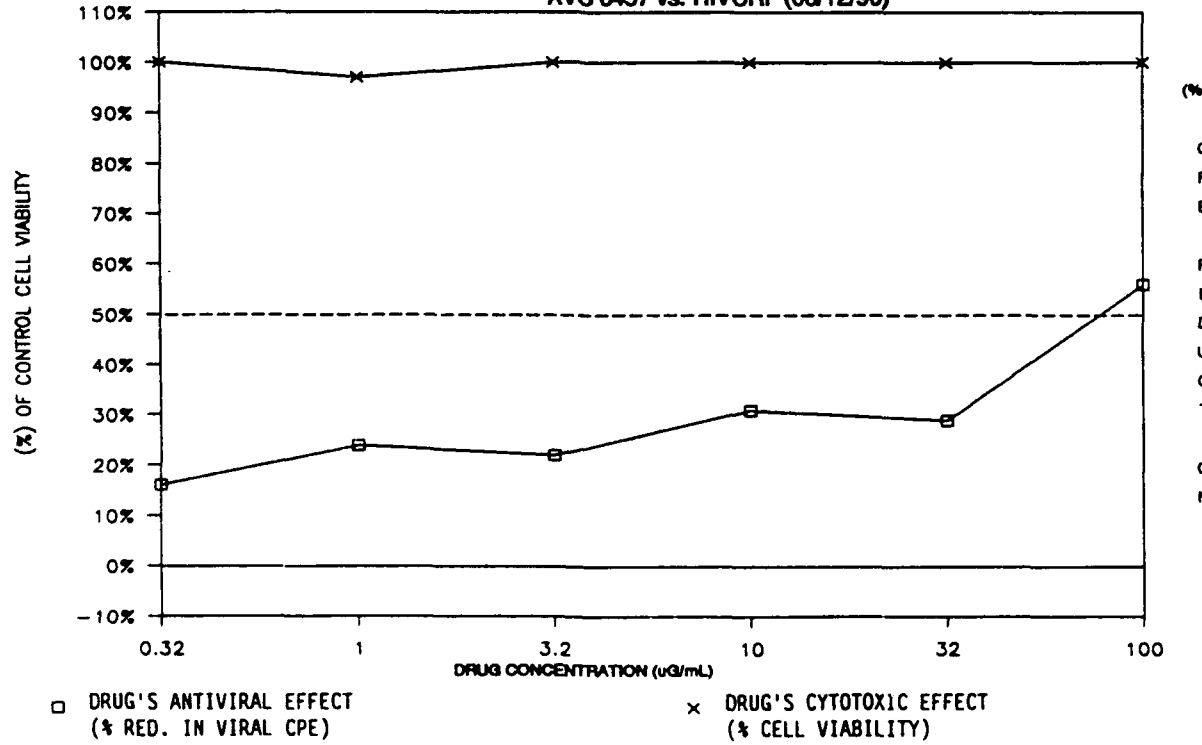
DRUG 6457		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/ml)	MEAN O.D.	% RED. IN VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	0.32	0.190	16%	1.473	100%	- .103
C	1	0.293	24%	1.307	97%	- .092
D	3.2	0.269	22%	1.442	100%	- .089
E	10	0.379	31%	1.462	100%	- .098
F	32	0.356	29%	1.436	100%	- .080
high G *	100	0.679	56%	1.406	100%	- .140

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6457 vs. HIVCRF (06/12/90)

□ DRUG'S ANTIVIRAL EFFECT
(% RED. IN VIRAL CPE)× DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

USAMRIID

Antiviral Drug Screening Program

09/21/90

STRUCTURE	SUBMITTER 01141.01	CTR NO MS-I-47	AVS NO AVS-006456
	DATE RECD 12-28-89	AMT RECEIVED [mg] 79.20	MOL WT (au) 544.694
HANDLING/STORAGE			
SOLUBILITY			
STABILITY			
ALT NAME			
SODIUM ETHYL(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE			

COMPOUND NAME

SODIUM ETHYL (CHOLESTERYLOXYCARBONYL) -PHOSPHONATE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

PLATE 1Q7
DRUG 6456IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6456
TAI: >10.33 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.463	0.349	0.539	0.566	0.386	0.388	0.253	0.256	0.399	0.448	0.239	0.199
B	1.590						tox		drug 6456 experimental		co/vc	tox
C	1.486						2.158	0.677	0.749	0.790	1.783	2.032
D	2.193						1.897	0.768	0.952	0.649	1.588	1.808
E	0.462						1.511	0.717	0.851	0.958	1.641	1.705
F	0.636						1.648	0.907	0.911	0.499	0.539	1.596
G	0.374						0.671	0.511	0.407	0.340	0.548	0.271
H							0.517	0.507	0.351	0.430	0.607	0.456
							0.437	0.442	0.455	0.454	0.455	0.522

toxic-cell toxicity

co-cell control

virus-virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

HIVCRF

PROJECT #

6520-2

CELLS

CEM Satisfactory

SPONSOR

USAIRRIID

SHIPMENT NUMBER

63

TEST DATE

06/12/90

STRN

RF2

DATE READ

06/19/90

REAGENT

0.449

	DRUG 6456	25%	50%	95%
VIRUS CONTROL	0.079	TC (uG/ml)	14.28	20.38
CELL CONTROL	1.265	IC (uG/ml)	2.39	-----
DIFFERENTIAL	1.186	ANTIVIRAL INDEX (AI)	5.92	-----

ROW ON PLATE	CONC. (uG/ml)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% RED. IN CPE	MEAN O.D.	% CELL VIABILITY	
Tow B	0.32	0.137	12%	1.573	100%	0.074
C	1	0.255	22%	1.397	100%	0.007
D	3.2	0.308	26%	1.154	91%	0.006
E	10	0.238	20%	1.167	92%	0.007
F	32	-.102	0%	0.029	2%	-.006
high G *	100	-.086	0%	0.050	4%	-.012

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6456 vs. HIVCRF (06/12/90)

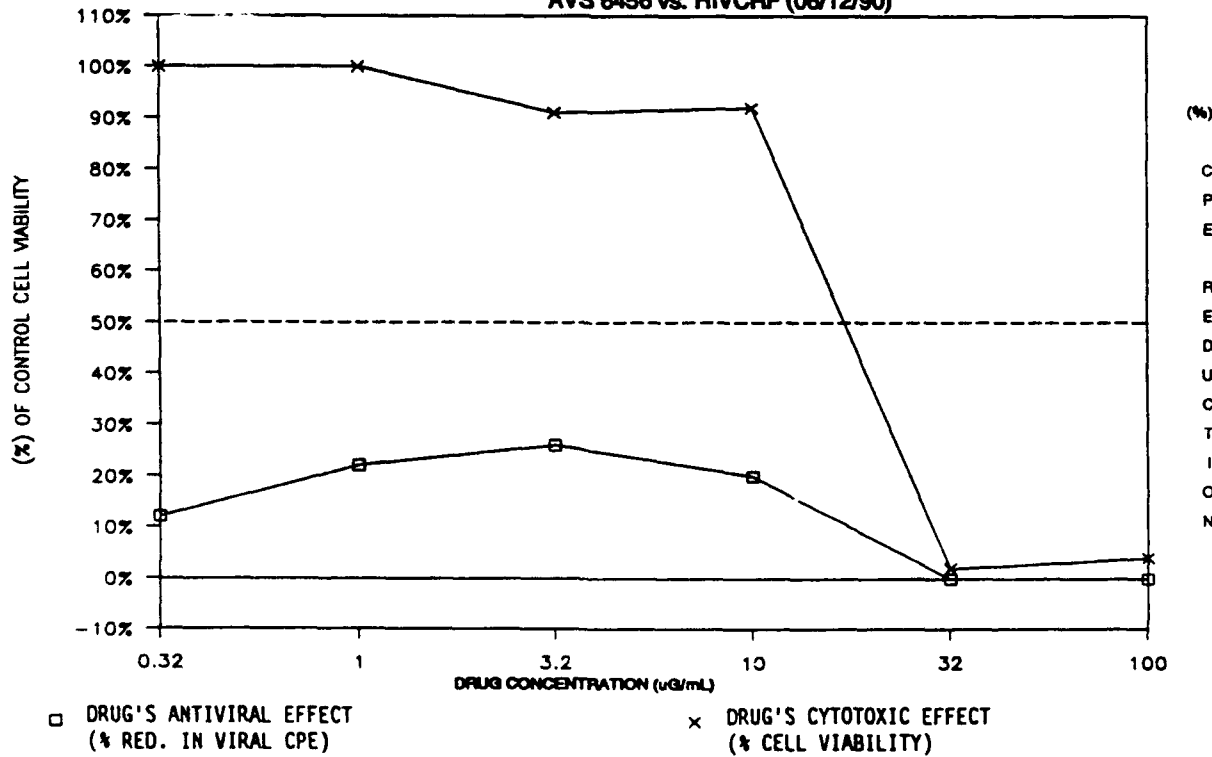


PLATE VCS
DRUG 6456IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6456
TAI: 0.00 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.064	0.056	0.057	0.057	0.059	0.058	0.002	0.001	0.002	0.002	0.002	0.002
B	1.137	1.382	0.224	0.255	0.236	1.253						
C	1.034	1.299	0.235	0.249	0.251	1.061						
D	0.984	1.077	0.361	0.349	0.350	0.932						
E	0.772	0.243	0.374	0.289	0.416	0.593						
F	0.044	0.242	0.035	0.035	0.034	0.037						
G	0.044	0.233	0.037	0.036	0.033	0.035						
H	0.040	0.049	0.051	0.051	0.051	0.053						

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug concn

values shown are optical densities

VIRUS	PT		PROJECT #	5975-1
CELLS	VERO	Satisfactory	SPONSOR	USAMRIID
SHIPMENT NUMBER	64	CONFIRMATORY #1 ** NEGATIVE **	TEST DATE	04/18/90
STRM	ADAMES		DATE READ	04/26/90
REAGENT	0.059	DRUG 6456	25%	50%
VIRUS CONTROL	0.185	TC (uG/mL)	43.30	120.00
CELL CONTROL	1.145	IC (uG/mL)	-----	-----
DIFFERENTIAL	0.961	ANTIVIRAL INDEX (AI)	-----	-----

ROW ON PLATE	CONC. (uG/mL)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
Tow B	3.2	0.001	100%	1.143	100%	- .006
C	10	0.010	99%	0.997	87%	- .008
D	32	0.118	88%	0.908	79%	- .008
E	100	0.124	87%	0.632	55%	- .008
F	320	.199	100%	-.008	0%	- .010
high G *	1000	-.189	100%	0.000	0%	- .019

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6456 vs. PT (04/18/90)

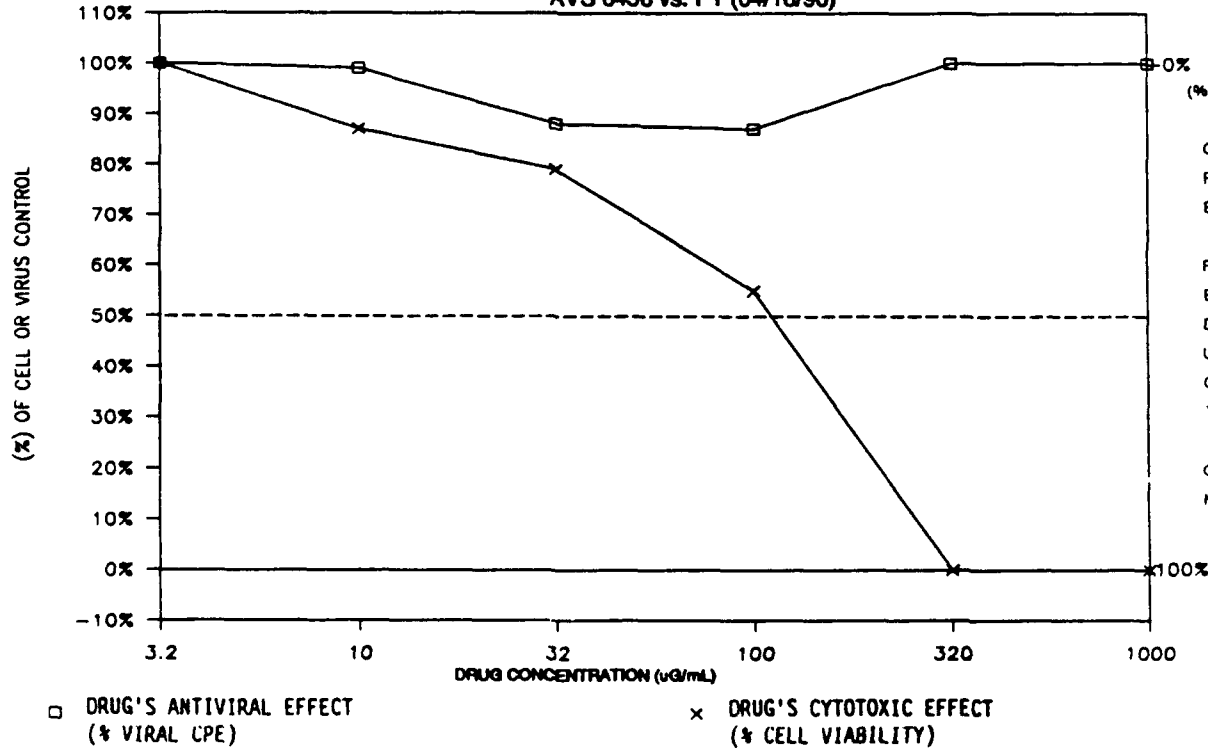


PLATE VCL
DRUG 6456IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6456
TAI: 0.00 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.056	0.052	0.051	0.051	0.052	0.052	0.001	0.002	0.001	0.001	0.001	0.001
B	0.906	1.126	0.044	0.043	0.040	1.015					0.781	
C	0.972	1.096	0.067	0.067	0.056	0.958					1.029	
D	1.066	1.047	0.077	0.071	0.061	1.002					0.946	
E	0.644	0.071	0.171	0.183	0.152	0.554					0.069	
F	0.041	0.095	0.038	0.039	0.037	0.038					0.092	
G	0.038	0.070	0.032	0.032	0.031	0.032					0.061	
H	0.038	0.047	0.048	0.053	0.049	0.048						

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc.

values shown are optical densities

VIRUS	SF	PROJECT #	5975-1
CELLS	VERO	SPONSOR	USAMRIID
SHIPMENT NUMBER	64	TEST DATE	04/18/90
STRN	SICILIAN	DATE READ	04/26/90
REAGENT	0.052	DRUG 6456	25%
VIRUS CONTROL	0.024	TC (µg/ml)	72.50
CELL CONTROL	0.952	IC (µg/ml)	130.00
DIFFERENTIAL	0.928	ANTIVIRAL INDEX (AI)	341.00

ROW ON PLATE	CONC. (µg/ml)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	3.2	-.030	100%	0.912	96%	-.004
C	10	-.010	100%	0.916	96%	-.003
D	32	-.008	100%	0.981	100%	0.001
E	100	0.096	90%	0.551	58%	-.004
F	320	-.033	100%	-.008	0%	-.005
high G *	1000	-.031	100%	-.003	0%	-.014

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6456 vs. SF (04/18/90)

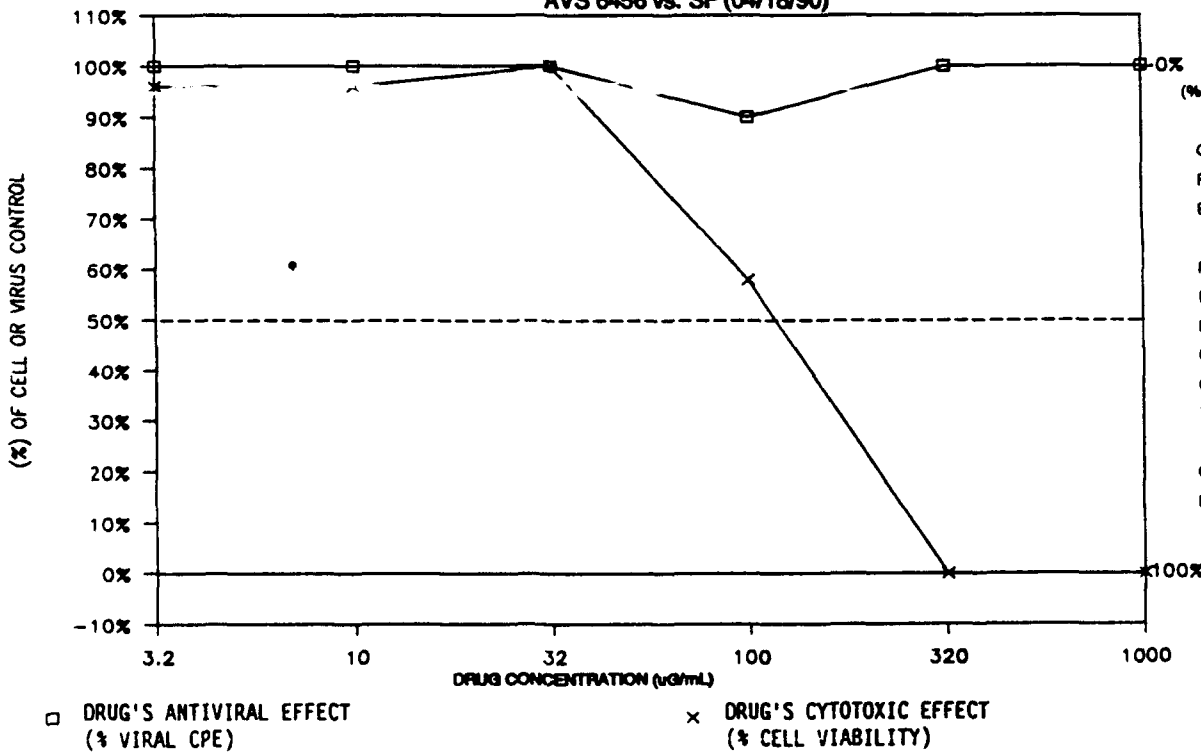


PLATE VBP
DRUG 6456

IN VITRO ANTIVIRAL RESULTS MTT ASSAY

DRUG: AVS 6456
TAI: 0.00 SI: _____

	1	2	3	4	5	6	7	8	9	10	11	12
A	reagent background						plastic background					
	0.086	0.068	0.066	0.068	0.071	0.066	0.001	0.002	0.002	0.001	0.002	0.002
B	1.336	1.491	0.112	0.091	0.111	1.375					1.479	
C	1.312	1.515	0.105	0.115	0.088	1.364					1.465	
D	1.290	1.521	0.083	0.086	0.099	1.282					1.444	
E	1.221	0.117	0.096	0.091	0.093	1.149					0.115	
F	0.410	0.126	0.138	0.196	0.147	0.378					0.120	
G	0.107	0.098	0.092	0.087	0.092	0.095					0.098	
H	drug 6456 colorimetric background											
H	0.057	0.073	0.064	0.060	0.062	0.065						

10³-cell toxicity control vs-virus control

BOLD = Highest drug score

values shown are optical densities.

VIRUS
CELLS
SHIPMENT NUMBER
STRN
REAGENT
VIRUS CONTROL
CELL CONTROL
DIFFERENTIAL

VE
 VERO Satisfactory
 64 CONFIRMATORY #1 ** NEGATIVE **
 TRINIDAD

0.071	DRUG 6466	25
0.042	TC { μ G/mL}	1
1.415	IC { μ G/mL}	
1.374	ANTIVIRAL INDEX (AI)	

PROJECT #	5975-1
SPONSOR	USAMRIID
TEST DATE	04/20/90
DATE READ	04/24/90

DRUG: 6455	25%	50%	95%
TC (uG/mL)	116.00	214.00	932.00
IC (uG/mL)	-----	-----	-----
ANTIVIRAL INDEX (AI)	-----	-----	-----

DRUG 6456		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (μ G/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY		
low B	3.2	- .002	100%	1.291	91%		-.006
C	10	- .001	100%	1.276	90%		-.009
D	32	- .012	100%	1.226	87%		-.011
E	100	- .012	100%	1.121	79%		-.007
F	320	0.046	97%	0.321	23%		0.002
high G *	1000	- .008	100%	0.044	3%		-.014

higher drug concentration test

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6456 vs. VE (04/20/90)

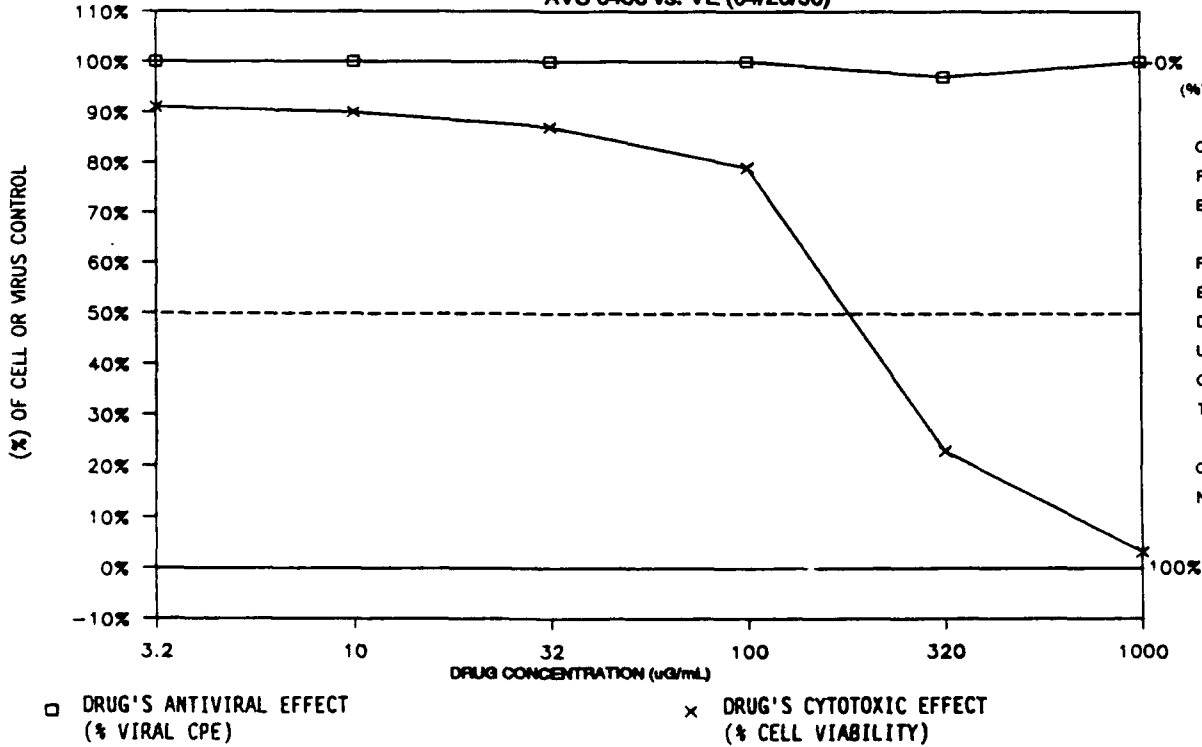


PLATE VAT
DRUG 6456IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6456
TAI: >15.51 SI: 2.16

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.074	0.076	0.073	0.073	0.073	0.074	0.001	0.001	0.000	0.001	0.001	0.001
B	0.976	1.014	0.301	0.302	0.311	0.917					0.995	
C	0.951	0.907	0.359	0.337	0.348	1.032					0.973	
D	0.954	0.913	0.509	0.517	0.494	0.966					0.989	
E	0.880	0.286	0.761	0.689	0.684	0.802					0.287	
F	0.117	0.292	0.097	0.082	0.079	0.080					0.294	
G	0.056	0.281	0.053	0.051	0.050	0.047					0.287	
H	0.052	0.065	0.069	0.069	0.069	0.068						

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS YF
 CELLS VERO Satisfactory; Active; Retest
 SHIPMENT NUMBER 64 CONFIRMATORY #1 ** GOOD **
 STRN ASIBI
 REAGENT 0.074

PROJECT # 5975-1

SPONSOR USAMRIID

TEST DATE 04/18/90

DATE READ 04/24/90

VIRUS CONTROL	0.214	DRUG 6456	25%	50%	95%
CELL CONTROL	0.891	TC (µg/ml)	132.00	198.00	317.00
DIFFERENTIAL	0.677	IC (µg/ml)	21.40	61.00	-----
		ANTIVIRAL INDEX (AI)	5.17	3.25	-----

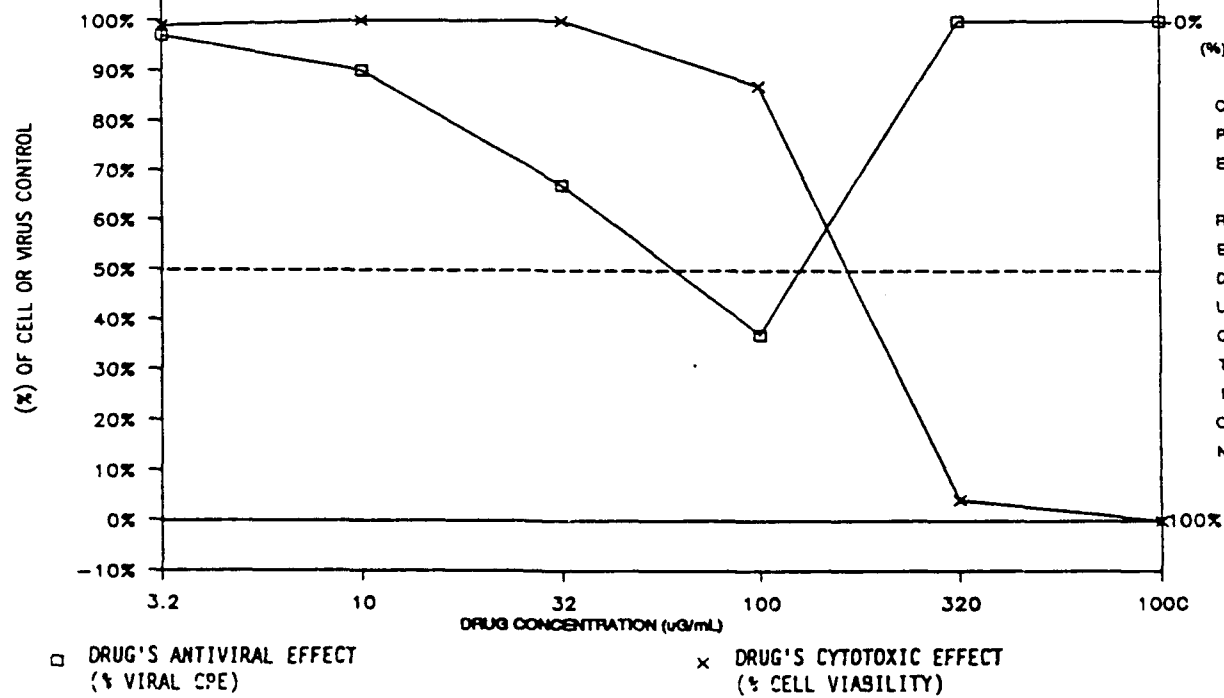
ROW ON PLATE	CONC. (µG/ml)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	3.2	0.023	97%	0.879	99%	- .006
C	10	0.065	90%	0.923	100%	- .005
D	32	0.224	67%	0.891	100%	- .005
E	100	0.429	37%	0.772	87%	- .005
F	320	-.193	100%	0.034	4%	- .009
high G *	1000	-.215	100%	0.000	0%	- .022

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6456 vs. YF (04/18/90)



USAMRIID

Antiviral Drug Screening Program

09/21/90

STRUCTURE	SUBMITTER 01141.01	CTR NO MS-I-44	AVS NO AVS-006455
	DATE RECD 12-28-89	AMT RECEIVED (mg) 79.20	MOL WT (au) 550.766
HANDLING/STORAGE			
SOLUBILITY			
STABILITY			
ALT NAME	DIETHYL(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE		

COMPOUND NAME

DIETHYL(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]						
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST	VH	RTE	LD50	MTC	LAB PR	DATE

PLATE 1Q7
DRUG 6455IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6455
TAI: >38.98 SI: >5.71

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.463	0.349	0.539	0.556	0.395	0.388	0.253	0.256	0.399	0.448	0.239	0.199
B	1.916	1.590	0.861	0.677	0.813	1.889					0.606	
C	1.840	1.486	0.820	0.986	0.873	2.017					1.588	
D	1.800	2.193	0.764	0.557	0.841	1.849					1.641	
E	2.057	0.462	0.856	0.811	1.334	2.016					0.539	
F	1.959	0.636	0.944	1.071	1.242	1.830					0.548	
G	2.003	0.374	1.269	1.478	1.180	1.913					0.607	
H	0.328	0.243	0.480	0.459	0.455	0.448						

tox=cell toxicity

ccv=cell control

vv=virus control

BOLD = highest drug concn

values shown are optical densities

VIRUS	HIVCRF	PROJECT #	6520-2
CELLS	CEM	SPONSOR	USAIRRIID
SHIPMENT NUMBER	63	TEST DATE	06/12/90
STRN	RF2	DATE READ	06/19/90
REAGENT	0.449	DRUG 6455	25%
VIRUS CONTROL	0.079	TC (uG/mL)	> 100.00
CELL CONTROL	1.265	IC (uG/mL)	0.49
DIFFERENTIAL	1.186	ANTIVIRAL INDEX (AI)	> 203.84

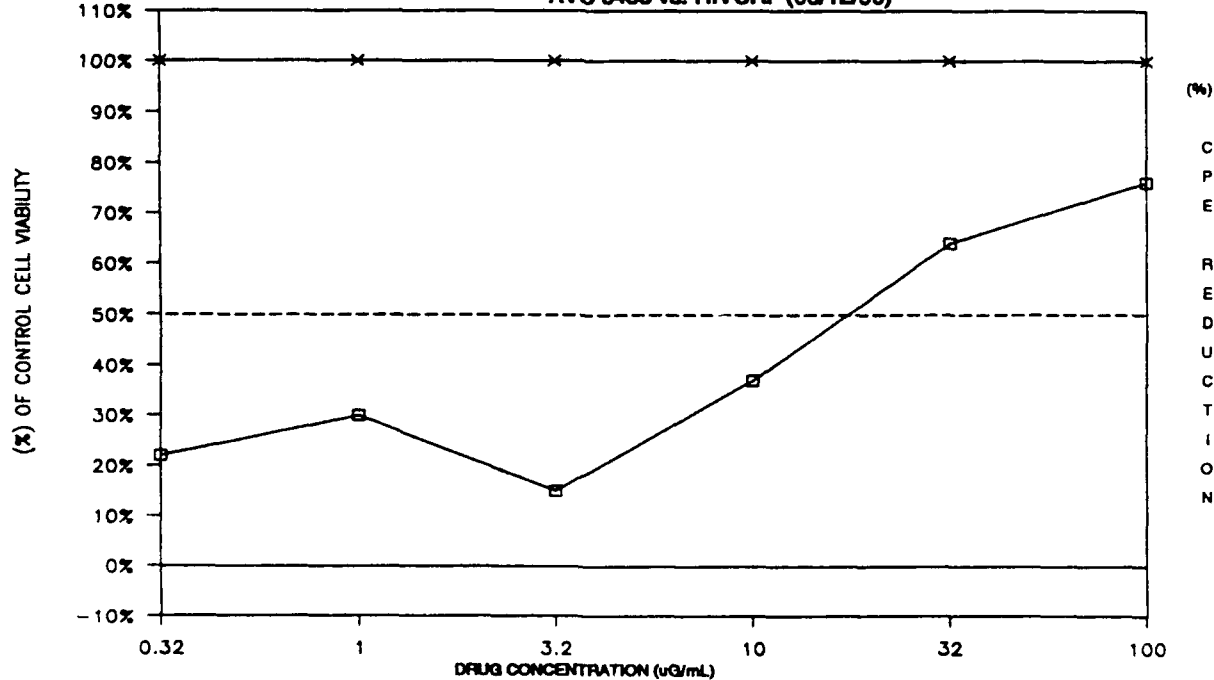
DRUG 6455		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/ml)	MEAN O.D.	% RED. IN VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	0.32	0.256	22%	1.454	100%	0.000
C	1	0.358	30%	1.473	100%	0.007
D	3.2	0.182	15%	1.365	100%	0.011
E	10	0.442	37%	1.557	100%	0.031
F	32	0.764	64%	1.652	100%	-.206
high G *	100	0.902	76%	1.631	100%	-.120

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6455 vs. HIVCRF (06/12/90)



DRUG'S ANTIVIRAL EFFECT
(% RED. IN VIRAL CPE)

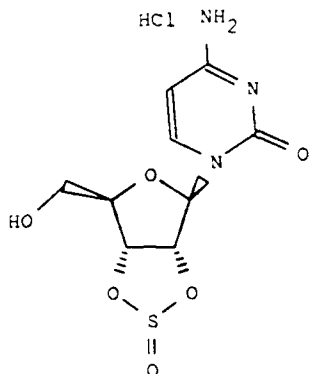
DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-II-71	AVS NO AVS-006462
		DATE RECD 12-28-89	AMT RECEIVED [mg] 72.40	MOL WT (au) 325.729
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME 2',3'-O-SULFINYL CYTIDINE HYDROCHLORIDE				



COMPOUND NAME 2',3'-O-SULFINYL CYTIDINE HYDROCHLORIDE
--

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE
IN VITRO SCREEN [ug/ml]	
VIR VR VR+ ID50 CELL MTC TI TI+ LAB PRT DATE	VIR RST VR VR+ DOSE MTC VEH RTE D TOX SP L PR DATE
HIV NOT ACT MT2 .06 0 SO MTT	
HIV NOT ACT MT2 < .32 0 SO MTT	
JE NOT ACT VERO 22.4 0 SO MTT 90-03-06	
PT NOT ACT VERO 38.6 0 SO MTT 90-03-06	
SF NOT ACT VERO 21 0 SO MTT 90-03-06	
VEE NOT ACT VERO 9.73 0 SO MTT 90-03-09	
VV 1.72 VERO 16.7 14.12 SO MTT	
VV 3.28 VERO 25.7 15.66 SO MTT	
YF NOT ACT VERO 44.4 0 SO MTT 90-03-06	

PLATE 055
DRUG 6462

IN VITRO ANTIVIRAL RESULTS MTT ASSAY

**DRUG: AVS 6462
TAI: >32.30 SI: 7.85**

	1	2	3	4	5	6	7	8	9	10	11	12
A	reagent background						plastic background					
	0.104	0.114	0.112	0.111	0.111	0.112	0.000	0.000	0.000	0.000	0.000	0.000
B	ca/va						tox	drug 6462 experimental			ca/va	tox
B	1.542						1.492	0.362	0.278	0.427	1.506	1.619
C	1.596						1.491	0.820	0.842	0.963	1.508	1.676
D	1.651						1.650	1.412	1.560	1.551	1.524	1.770
E	0.160						1.040	0.929	0.938	0.852	0.247	1.058
F	0.248						0.573	0.511	0.523	0.524	0.174	0.542
G	0.213						0.320	0.312	0.315	0.305	0.238	0.349
H							drug 6462 colorimetric background					
H							0.153	0.123	0.110	0.108	0.110	0.117

VIRUS	VV		PROJECT #	5975-4
CELLS	VERO	Satisfactory; Active; Retest	SPONSOR	USAMRIID
SHIPMENT NUMBER	63	RETEST AT 100 UG/ML	TEST DATE	03/29/90
STRN	LRDCA		DATE READ	04/04/90
REAGENT	0.111	DRUG 6462	25%	50%
VIRUS CONTROL	0.103	TC (uG/mL)	25.70	61.10 > 320.00
CELL CONTROL	1.444	IC (uG/mL)	1.56	3.28 9.54
DIFFERENTIAL	1.341	ANTIVIRAL INDEX (AI)	16.44	18.56 > 33.56

DRUG 6462		ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CC TROL
ROW ON PLATE	CONC. (μ G/mL)	MEAN O.D.	% VIRAL CPK	MEAN O.D.	% CELL VIABILITY			
low B	1	0.136	90%	1.439	100%			1.006
C	3.2	0.663	51%	1.474	100%			-0.001
D	10	1.297	3%	1.602	100%			-0.003
E	32	0.694	48%	0.939	65%			-0.001
F	100	0.294	78%	0.435	30%			0.012
high G	320	0.055	96%	0.182	13%			0.042

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

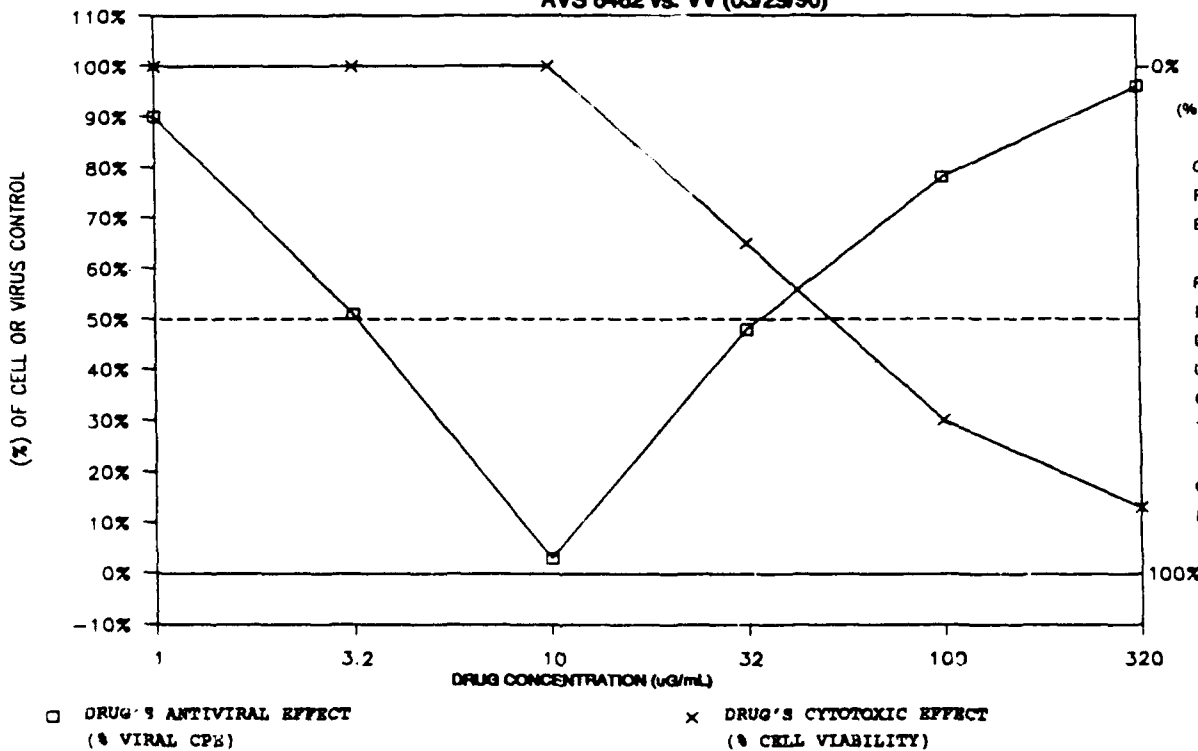


PLATE 0U2
DRUG 6462IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6462
TAI: 30.00 SI: 9.69

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.125	0.117	0.119	0.120	0.124	0.130	0.000	0.000	0.000	0.000	0.000	0.000
B	1.284	1.387	0.125	0.155	0.151	1.443						
C	1.197	1.494	0.453	0.332	0.473	1.315						
D	1.183	1.466	1.131	1.258	1.282	1.466						
E	1.311	0.176	1.234	1.216	1.347	1.456						
F	0.415	0.176	0.359	0.399	0.430	0.446						
G	0.280	0.197	0.263	0.256	0.255	0.294						
H	0.124	0.106	0.110	0.108	0.103	0.152						

low=cell toxicity

cc=cell control

vv=virus control

BOLD = highest drug conc.

values shown are optical densities

VIRUS

VV

PROJECT #

5975-4

CELLS

VERO

SPONSOR

USAMRIID

SHIPMENT NUMBER

Satisfactory

TEST DATE

04/19/90

STRM

LEDCA

DATE READ

04/25/90

REAGENT

DRUG 6462

25%

50%

95%

VIRUS CONTROL

TC (μg/ml)

16.70

24.40

> 100.00

CELL CONTROL

IC (μg/ml)

1.10

1.72

DIFFERENTIAL

ANTIVIRAL INDEX (AI)

15.27

14.12

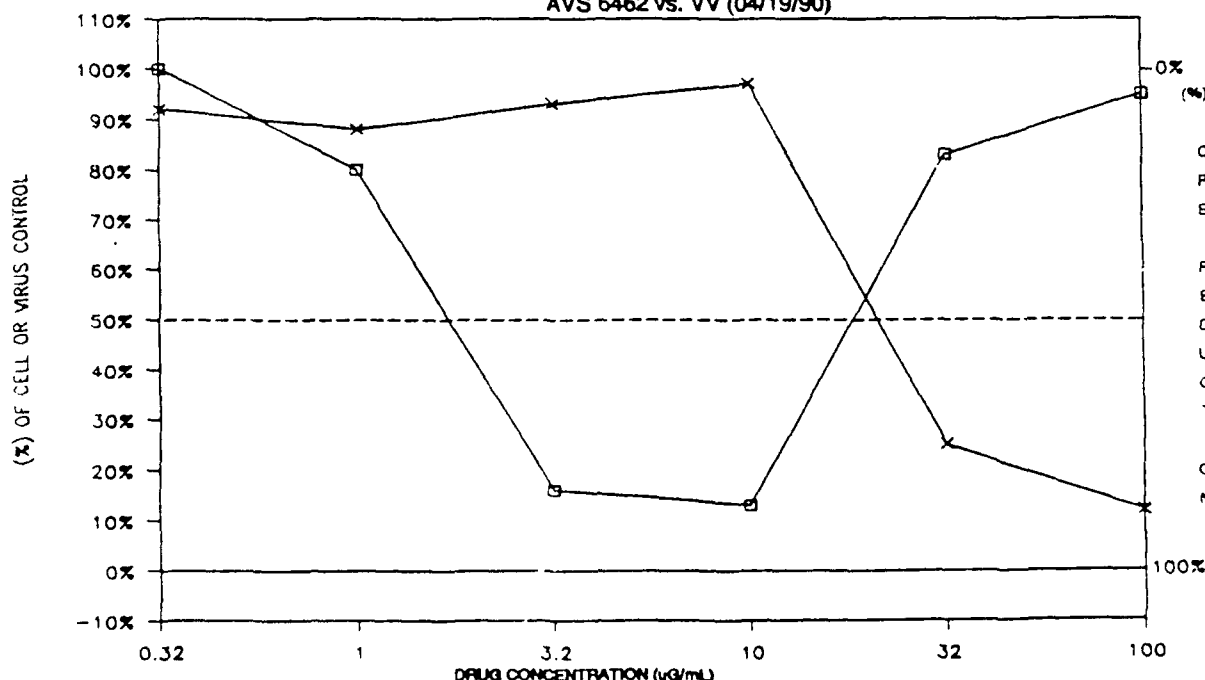
LOW ON PLATE	CONC. (μg/ml)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
Tow B	0.32	.084	100%	1.211	92%	0.029
C	1	0.242	80%	1.154	88%	-.020
D	3.2	1.041	16%	1.217	93%	-.015
E	10	1.081	13%	1.274	97%	-.013
F	32	0.215	83%	0.325	25%	-.017
high G *	100	0.059	95%	0.164	12%	0.001

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6462 vs. VV (04/19/90)



DRUG'S ANTIVIRAL EFFECT
(% VIRAL CPE)

DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

USAMRIID

Antiviral Drug Screening Program

05/18/91

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-II-95	AVS NO AVS-006467
		DATE RECD 12-28-89	AMT RECEIVED [mg] 72.60	MOL WT (au) 261.667
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME				
2',O2-Anhydrocytidine Hydrochloride				

COMPOUND NAME

2'-O²-ANHYDROCYTIDINE HYDROCHLORIDE

PLATE 0U4
DRUG 6467IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6467
TAI: 23.24 SI: 4.65

	1	2	3	4	5	6	7	8	9	10	11	12
A	0.105	0.097	0.123	0.114	0.109	0.119	0.000	0.000	0.000	0.000	0.000	0.000
B	1.314						1.488	0.153	0.166	0.134	1.321	1.330
C	1.339						1.510	0.179	0.148	0.205	1.470	1.424
D	1.312						1.553	0.258	0.206	0.325	1.410	1.427
E	0.182						1.415	1.153	1.398	1.302	0.160	1.541
F	0.180						1.058	0.847	0.883	0.733	0.173	0.825
G	0.174						0.450	0.426	0.446	0.427	0.142	0.400
H							0.107	0.110	0.105	0.108	0.114	0.117

tox=cell toxicity cc=cell control vv=virus control

BOLD = highest drug conc.

values shown are optical densities

VIRUS

VV

PROJECT #

5975-4

CELLS

VERO Satisfactory; Active; Retest

SPONSOR

USAMRIID

SHIPMENT NUMBER

63 RETEST AT 3.2 UG/ML

TEST DATE

04/19/90

STRN

LEDCA

DATE READ

04/25/90

REAGENT

0.111

DRUG 6467

25%

50%

95%

VIRUS CONTROL

0.057

TC (uG/mL)

0.82

1.86

> 3.20

CELL CONTROL

1.250

IC (uG/mL)

0.13

0.18

DIFFERENTIAL

1.193

ANTIVIRAL INDEX (AI)

6.52

10.53

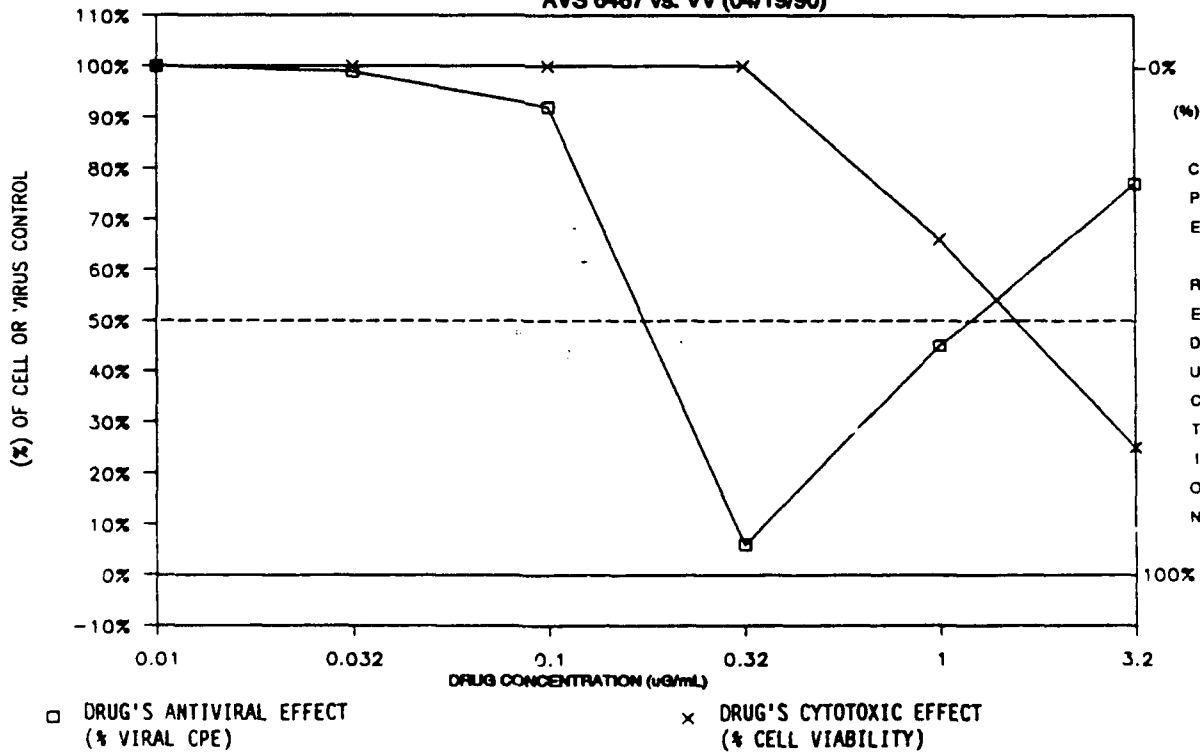
DRUG 6467		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
Tow B	0.01	-.023	100%	1.292	100%	0.006
C	0.032	0.006	99%	1.353	100%	0.003
D	0.1	0.098	92%	1.382	100%	-.003
E	0.32	1.122	6%	1.373	100%	-.006
F	1	0.654	45%	0.831	66%	-.001
high G *	3.2	0.269	77%	0.318	25%	-.004

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6467 vs. VV (04/19/90)



USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-V-109	AVS NO AVS-006443
		DATE RECD 12-28-89	AMT RECEIVED [mg] 86.00	MOL WT (au) 818.979
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME N3-BENZYL-2',5'-DI-O-TRITYLURIDINE				

COMPOUND NAME

N3-BENZYL-2',5'-DI-O-TRITYLURIDINE

PLATE UQ9

IN VITRO ANTIVIRAL RESULTS
MTT ASSAYDRUG: AVS 6443
TAI: >8.74 SI: 0.00

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.061	0.061	0.061	0.059	0.059	0.062	0.001	0.001	0.002	0.002	0.001	0.002
tox	oc/va		drug 6443 experimental		tox						oc/va	
B	1.421	1.475	0.417	0.536	0.571	1.230					1.590	
C	1.514	1.258	0.406	0.411	0.448	0.783					0.962	
D	1.327	1.605	0.658	0.571	0.499	1.230					1.398	
E	1.281	0.452	0.761	0.771	0.763	1.170					0.434	
F	1.222	0.408	0.872	0.841	0.865	1.158					0.366	
G	0.678	0.466	0.284	0.271	0.382	0.582					0.425	
H	0.207	0.089	0.068	0.064	0.064	0.064						
	tox-cell toxicity	con-cell control	vc-virus control				BOLD = highest drug conc					

values shown are optical densities

VIRUS	YF	PROJECT #	5975-1
CELLS	VERO	Sponsor	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	03/22/90
STRN	ASIBI	DATE READ	03/30/90
REAGENT	0.061		
		DRUG 6443	
VIRUS CONTROL	0.365		25%
CELL CONTROL	1.321		50%
DIFFERENTIAL	0.956		95%
		TC (uG/mL)	427.00
		IC (uG/mL)	56.60
		ANTIVIRAL INDEX (AI)	7.54

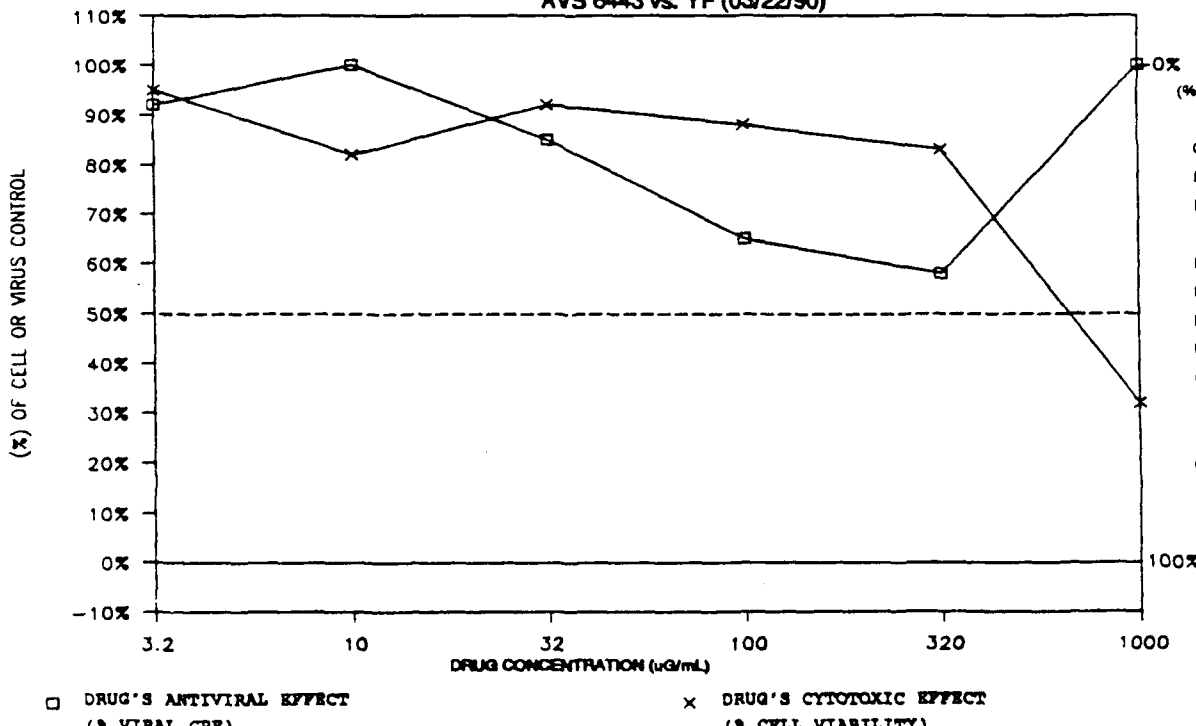
ROW ON PLATE	CONC. (uG/mL)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY		
low B	3.2	0.079	92%	1.261	95%		0.004
C	10	-0.08	100%	1.084	82%		0.004
D	32	0.147	85%	1.214	92%		0.004
E	100	0.332	65%	1.157	88%		0.008
F	320	0.405	58%	1.101	83%		0.029
high G *	1000	-.260	100%	0.423	32%		0.147

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6443 vs. YF (03/22/90)



USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-VII-83	AVS NO AVS-006444
		DATE RECD 12-28-89	AMT RECEIVED [mg] 79.00	MOL WT (au) 726.837
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME				
3'-DEOXY-2',5'-DI-O-TRITYL-3'-OXOURIDINE				

COMPOUND NAME

3'-DEOXY-2',5'-DI-O-TRITYL-3'-OXOURIDINE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV> AD2 >VSV	HOST VH RTE LD50 MTC LAB PR DATE

PLATE UQO
DRUG 6444

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6444
TAI: 3.21 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background											
A	0.077	0.073	0.071	0.070	0.072	0.072	0.001	0.001	0.002	0.001	0.002	0.002
B		cell					tox	drug 6444 experimental	cell	tox		
C		1.519					1.517	0.509	0.525	0.554	1.544	1.671
D		1.478					1.483	0.458	0.481	0.478	1.484	1.577
E		1.487					1.474	0.462	0.467	0.492	1.537	1.384
F		0.607					1.542	0.621	0.651	0.640	0.589	1.615
G		0.615					1.327	0.845	0.897	0.927	0.619	1.517
H		0.626					0.697	0.474	0.515	0.524	0.613	0.680
	drug 6444 colorimetric background											
							0.121	0.082	0.081	0.078	0.078	0.077

tox=cell toxicity cc=cell control vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

JE

CELLS

VERO Satisfactory; Active; Retest

SHIPMENT NUMBER

63 TOXICITY RERUN

STRM

NARAYAMA

PROJECT # 5975-1

SPONSOR USAMRIID

TEST DATE 03/22/90

DATE READ 03/29/90

REAGENT

0.073 DRUG 6444

25%

50%

95%

VIRUS CONTROL

0.539 TC (uG/mL)

547.00

861.00

> 1000.00

CELL CONTROL

1.436 IC (uG/mL)

260.00

DIFFERENTIAL

0.897 ANTIVIRAL INDEX (AI)

2.10

DRUG 6444		ANTIVIRAL TEST VALUES			CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	VIABILITY		
low B	3.2	-0.087	100%	1.517	100%	100%	0.005	
C	10	-0.145	100%	1.452	100%	94%	0.006	
D	32	-0.144	100%	1.351	100%	94%	0.006	
E	100	0.017	98%	1.497	100%	100%	0.009	
F	320	0.268	70%	1.340	93%	93%	0.010	
high G	1000	-0.156	100%	0.567	39%	39%	0.049	

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6444 vs. JE (03/22/90)

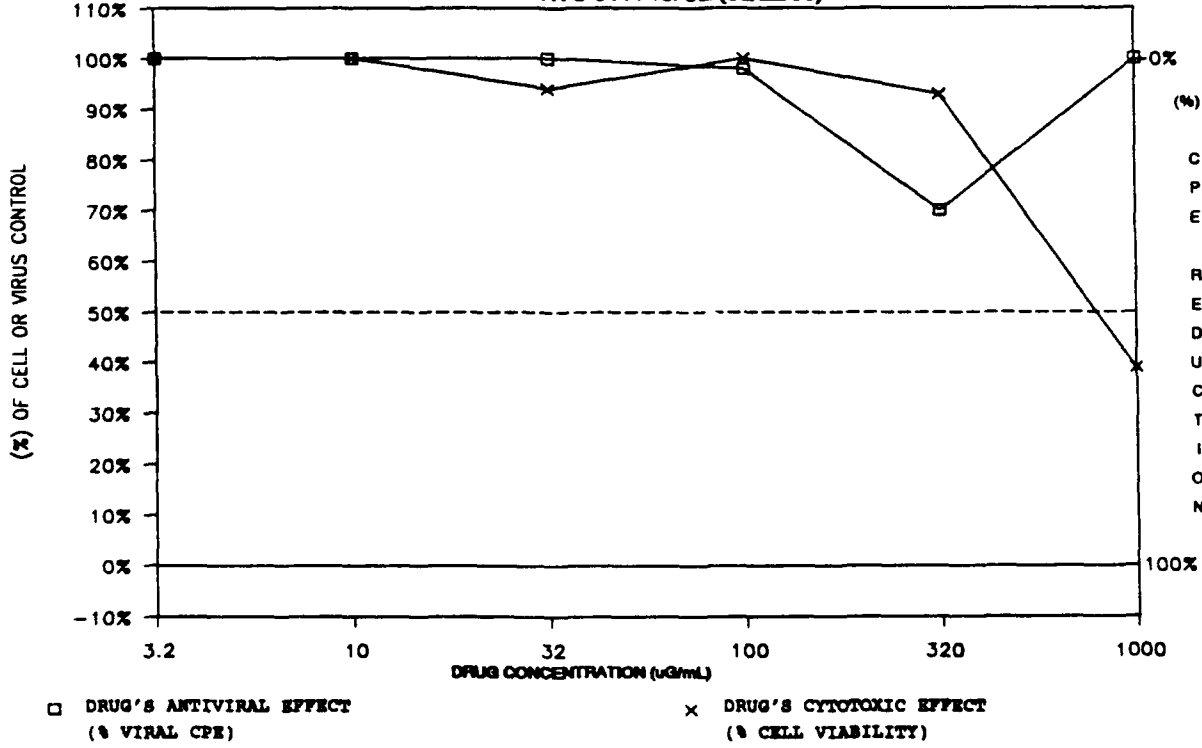


PLATE URI
DRUG 6444

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6444
TAI: 3.44 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
reagent background												
A	0.054	0.055	0.054	0.051	0.051	0.053	0.002	0.002	0.001	0.001	0.001	0.001
B		1.408					1.0x	drug 6444 experimental	cc/vc	tox		
C		1.631					1.654	0.253	0.349	0.342	1.305	1.302
D		1.634					1.553	0.377	0.414	0.423	1.539	1.238
E		0.388					1.452	0.477	0.492	0.519	1.485	1.209
F		0.391					1.415	0.593	0.743	0.580	0.418	1.296
G		0.316					1.118	0.988	0.750	0.778	0.386	1.001
H							0.384	0.422	0.430	0.447	0.403	0.380
							drug 6444 colorimetric background					
							0.055	0.057	0.056	0.055	0.051	0.049

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

PT

PROJECT # 5975-1

CELLS

VERO

Satisfactory; Active; Retest

SPONSOR USAMRIID

SHIPMENT NUMBER

63 TOXICITY RERUN

TEST DATE 03/22/90

STRN

ADAMES

DATE READ 03/30/90

REAGENT

DRUG 6444

25%

50%

95%

VIRUS CONTROL

TC (uG/mL)

257.00

601.00

> 1000.00

CELL CONTROL

IC (uG/mL)

115.00

DIFFERENTIAL

ANTIVIRAL INDEX (AI)

2.24

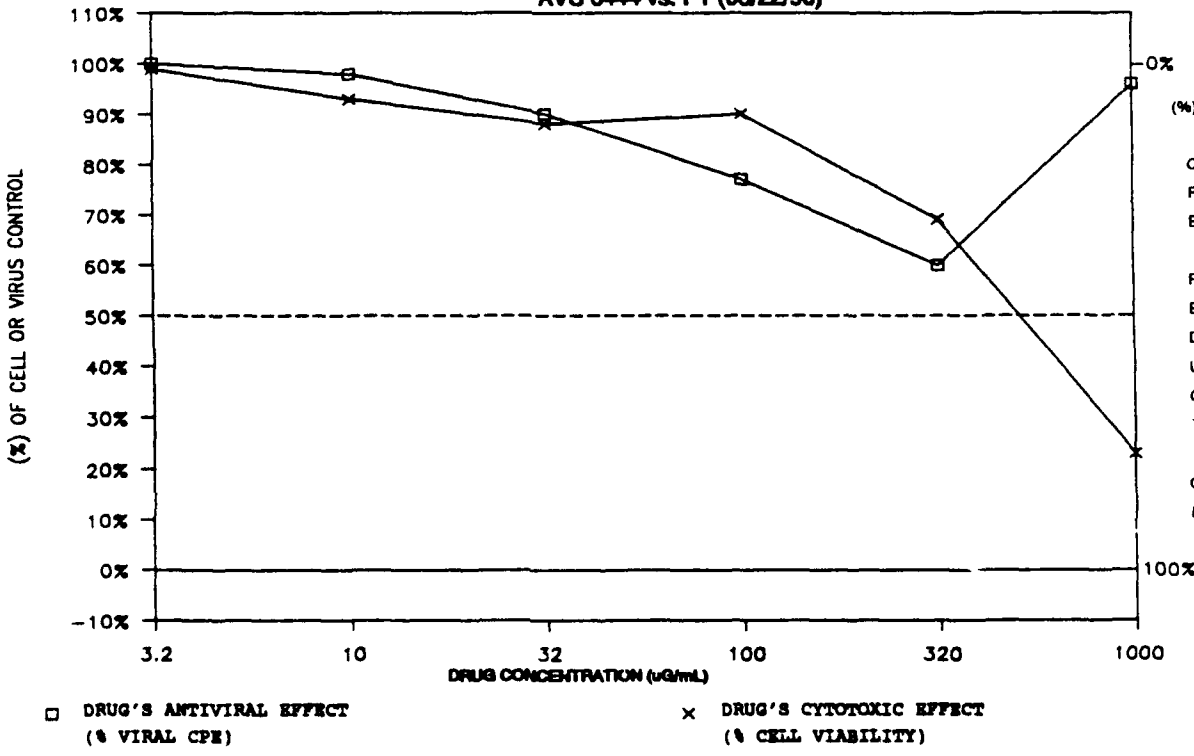
DRUG 6444		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	3.2	-0.065	100%	1.429	99%	-0.004
C	10	0.023	98%	1.345	93%	-0.002
D	32	0.110	90%	1.276	88%	0.002
E	100	0.252	77%	1.300	90%	0.003
F	320	0.451	60%	1.003	69%	0.004
high G *	1000	0.047	96%	0.327	23%	0.002

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6444 vs. PT (03/22/90)



USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE	CHIRAL	SUBMITTER	CTR NO	AVS NO
		01141.01	KN-VII-21	AVS-006445
		DATE RECD 12-28-89	AMT RECEIVED [mg] 74.00	MOL WT (au) 726.837
		HANDLING/STORAGE		
		SOLUBILITY		
		STABILITY		
		ALT NAME 2'-DEOXY-3',5'-DI-O-TRITYL-2'-OXOURIDINE		

COMPOUND NAME

2'-DEOXY-3',5'-DI-O-TRITYL-2'-OXOURIDINE

PLATE UQA
DRUG 6445

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6445
TAI: >18.33 SI: 3.27

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background						plastic background					
A	0.063	0.063	0.062	0.060	0.056	0.061	0.001	0.001	0.001	0.001	0.001	0.001
B	1.549	1.409	0.619	0.596	0.557	1.504					1.418	
C	1.442	1.368	0.740	0.805	0.674	1.607					1.392	
D	1.368	1.448	0.998	0.971	0.924	1.488					1.550	
E	1.067	0.526	0.934	0.600	0.848	1.022					0.433	
F	0.062	0.481	0.051	0.050	0.054	0.051					0.380	
G	0.066	0.494	0.066	0.061	0.059	0.065					0.400	
	drug 6445 colorimetric background											
H	0.075	0.071	0.065	0.062	0.064	0.062						

TOX=cell toxicity CC=cell control VC=virus control

BOLD = highest drug conc.

values shown are optical densities

VIRUS	YF	PROJECT #	5975-1
CELLS	VERO	Sponsor	USAMRIID
SHIPMENT NUMBER	63	TEST DATE	03/22/90
STRN	AS IBI	DATE READ	03/30/90
REAGENT	0.061	DRUG 6445	25%
VIRUS CONTROL	0.392	TC (uG/mL)	29.60
CELL CONTROL	1.370	IC (uG/mL)	2.35
DIFFERENTIAL	0.979	ANTIVIRAL INDEX (AI)	12.63
			50%
			95%

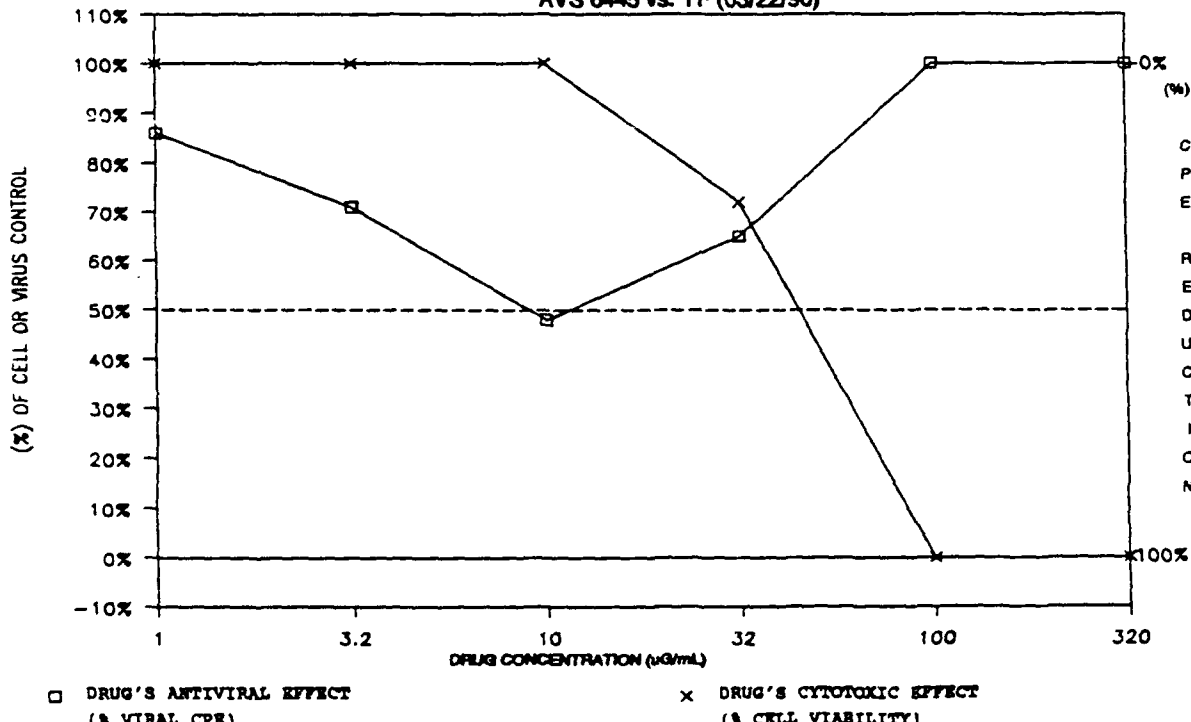
DRUG 6445		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (uG/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	1	0.137	86%	1.465	100%	0.001
C	3.2	0.284	71%	1.461	100%	0.003
D	10	0.511	48%	1.166	100%	0.001
E	32	0.338	65%	0.980	72%	0.004
F	100	-0.411	100%	-0.014	0%	0.010
high G	320	-0.404	100%	-0.009	0%	0.014

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6445 vs. YF (03/22/90)



USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE	CHIRAL	SUBMITTER 01141.01	CTR NO KN-II-53	AVS NO AVS-006449
		DATE RECD 12-28-89	AMT RECEIVED [mg] 75.00	MOL WT (au) 284.271
HANDLING/STORAGE				
SOLUBILITY				
STABILITY				
ALT NAME 2',3'-O-ISOPROPYLIDINEURIDINE				

COMPOUND NAME

2', 3'-O-ISOPROPYLIDINEURIDINE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

USAMRIID

Antiviral Drug Screening Program

25/18/90

STRUCTURE	SUBMITTER 01141.01	CTR NO MS-I-47	AVS NO AVS-006456
	DATE RECD 12-28-89	AMT RECEIVED [mg] 79.20	MOL WT (au) 544.694
	HANDLING/STORAGE		
	SOLUBILITY		
	STABILITY		
	ALT NAME SODIUM ETHYL(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE		

COMPOUND NAME

SODIUM ETHYL (CHOLESTERYLOXYCARBONYL) -PHOSPHONATE

PLATE UDN
DRUG 6456

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6456
TAI: 3.19 SI: _____

	1	2	3	4	5	6	7	8	9	10	11	12
reagent background												
A	0.059	0.057	0.063	0.055	0.063	0.054	0.001	0.001	0.002	0.002	0.001	0.002
B	0.936	0.768	0.225	0.228	0.213	0.841					0.941	
C	0.893	0.819	0.172	0.162	0.145	0.951					1.024	
D	0.989	0.750	0.144	0.136	0.135	1.052					0.789	
E	0.964	0.281	0.432	0.408	0.423	0.955					0.268	
F	0.174	0.271	0.053	0.051	0.052	0.077					0.265	
G	0.050	0.280	0.046	0.044	0.044	0.043					0.283	
drug 6456 colorimetric background												
H	0.050	0.053	0.055	0.057	0.052	0.055						

tox=cell toxicity cc=cell control vc=virus control

BOLD = highest drug concn

values shown are optical densities

VIRUS

SF

CELLS

VERO Satisfactory; Active; Retest

PROJECT #

5975-1

SHIPMENT NUMBER

63

SPONSOR

USAMRIID

STRM

SICILIAN

TEST DATE

03/06/90

REAGENT

0.059

DATE READ

03/14/90

VIRUS CONTROL

0.216

	DRUG 6456	25%	50%	95%
TC (ug/mL)		50.70	69.40	198.00
IC (ug/mL)		30.60	-----	-----
ANTIVIRAL INDEX (AI)		1.66	-----	-----

CELL CONTROL

0.790

DIFFERENTIAL

0.574

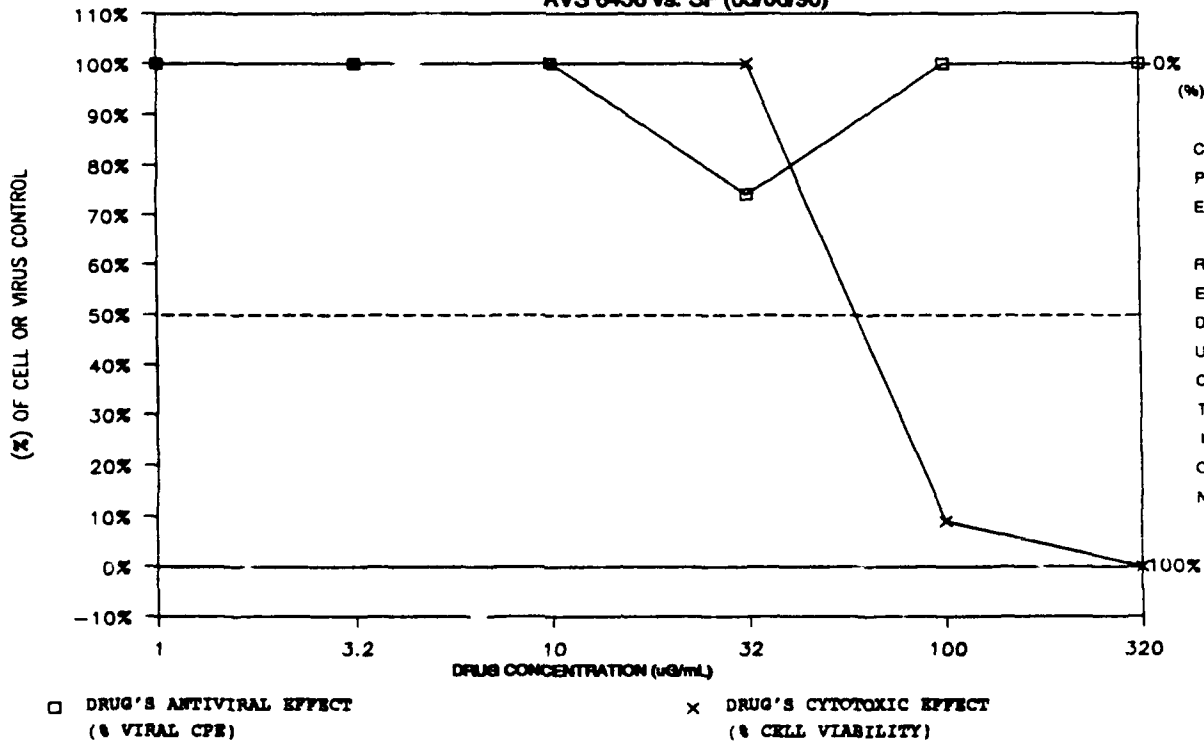
DRUG 6456		ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES			COLORIMETRIC CONTROL
ROW ON PLATE	CONC. (ug/mL)	MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY		
low B	1	-.049	100%	0.834	100%		-.004
C	3.2	-.108	100%	0.871	100%		-.007
D	10	-.135	100%	0.963	100%		-.001
E	32	0.150	74%	0.905	100%		-.004
F	100	-.217	100%	0.073	9%		-.006
high G *	320	-.222	100%	-.004	0%		-.008

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6456 vs. SF (03/06/90)

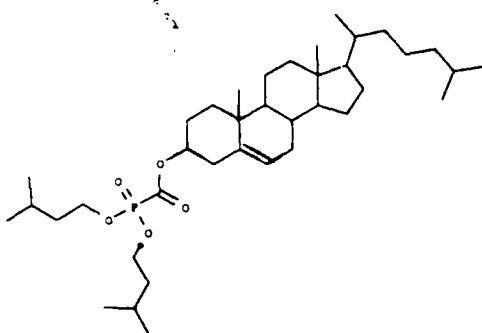


USAMRIID

Antiviral Drug Screening Program

05/18/90

STRUCTURE



SUBMITTER 01141.01	CTR NO KN-I-105	AVS NO AVS-006458
DATE RECD 12-28-89	AMT RECEIVED [mg] 70.40	MOL WT (au) 634.929
HANDLING/STORAGE		
SOLUBILITY		
STABILITY		
ALT NAME DI-ISOBUTYL(CHOLESTERYLOXYCARBONYL)-PHOSPHONATE		

COMPOUND NAME

DI-ISOBUTYL (CHOLESTERYLOXYCARBONYL)-PHOSPHONATE

SCREEN INSTRUCTION	IN VIVO TOXICITY [mg/kg]
PRIORITY=PT>VEE>YF>KHF>PIC>JE>SF>VV>AD2>VSV	HOST VH RTE LD50 MTC LAB PR DATE

PLATE UCO

IN VITRO ANTIVIRAL RESULTS
MTT ASSAY

DRUG: AVS 6458

DRUG 6458

TAI: >0.50 SI: -----

	1	2	3	4	5	6	7	8	9	10	11	12
	reagent background											
A	0.062	0.059	0.057	0.058	0.057	0.058	0.002	0.001	0.001	0.002	0.001	0.001
	tox	curve	drug 6458 experimental		tox						curve	
B	1.334	1.245	0.411	0.439	0.415	1.113					1.215	
C	1.259	1.243	0.405	0.394	0.392	1.110					1.185	
D	1.298	1.283	0.365	0.422	0.412	1.110					1.211	
E	1.203	0.413	0.454	0.375	0.451	1.054					0.442	
F	1.226	0.403	0.443	0.430	0.497	1.030					0.455	
G	1.024	0.393	0.689	0.669	0.689	0.862					0.446	
	drug 6458 colorimetric background											
H	0.058	0.061	0.061	0.060	0.060	0.061						

tox=cell toxicity

cc=cell control

vc=virus control

BOLD = highest drug conc

values shown are optical densities

VIRUS

CELLS

SHIPMENT NUMBER

STRM

REAGENT

VIRUS CONTROL

CELL CONTROL

DIFFERENTIAL

PT

VERO

Satisfactory; Active; Rerest

PROJECT #

5975-1

SPONSOR

USAMRIID

TEST DATE

03/06/90

DATE READ

03/16/90

	DRUG 6458	25%	50%	95%
TC (uG/mL)	> 320.00	> 320.00	> 320.00	
IC (uG/mL)	239.00	-----	-----	
ANTIVIRAL INDEX (AI)	> 1.34	-----	-----	

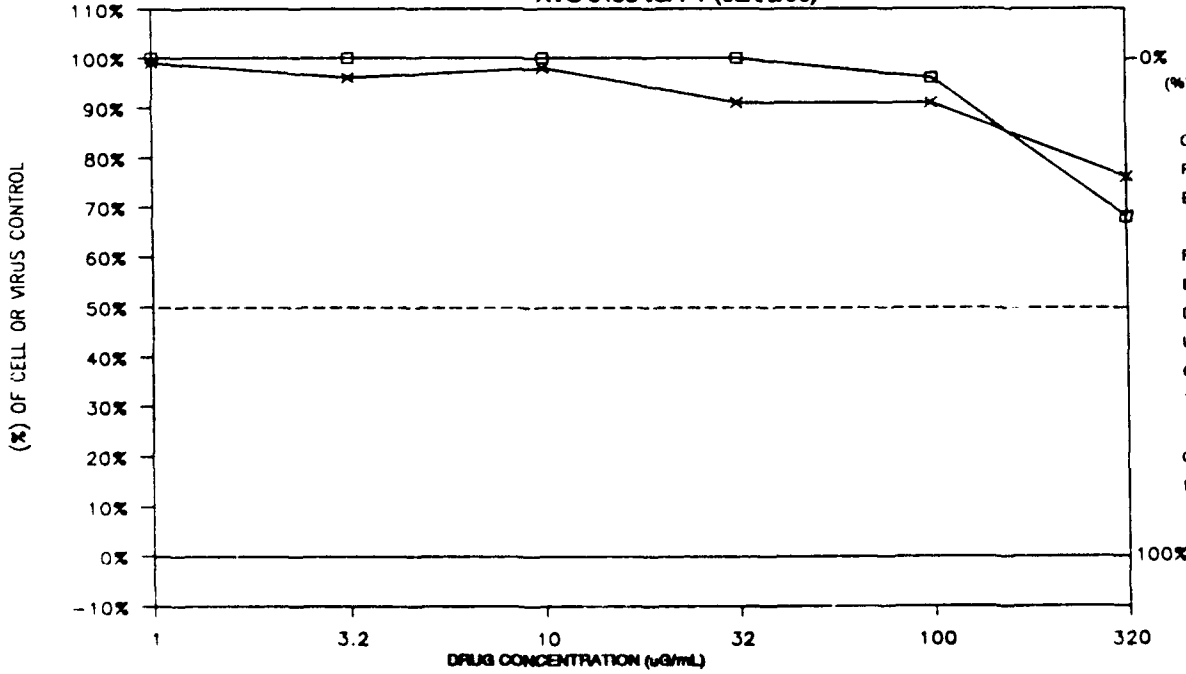
ROW ON PLATE	CONC. (uG/mL)	ANTIVIRAL TEST VALUES		CYTOTOXICITY TEST VALUES		COLORIMETRIC CONTROL
		MEAN O.D.	% VIRAL CPE	MEAN O.D.	% CELL VIABILITY	
low B	1	-.006	100%	1.163	99%	0.002
C	3.2	-.029	100%	1.125	96%	0.001
D	10	-.027	100%	1.145	98%	0.001
E	32	-.001	100%	1.068	91%	0.002
F	100	0.029	95%	1.068	91%	0.002
high G *	320	0.257	68%	0.885	76%	0.000

* highest drug concentration tested

values shown are final adjusted numbers

SUMMARY GRAPH

AVS 6458 vs. PT (03/06/90)



DRUG'S ANTIVIRAL EFFECT
(% VIRAL CPE)

DRUG'S CYTOTOXIC EFFECT
(% CELL VIABILITY)

APPENDIX V

SUMMARY OF COMPOUNDS SHOWING SIGNIFICANT ANTIVIRAL ACTIVITY.

The table summarizes those compounds shown to have significant antiviral activity against one or more of the test viruses. The test data together with the I_{50} are given in the sheets of Appendix 3.

AVS #S OF COMPOUNDS SHOWING SIGNIFICANT ANTI-VIRAL ACTIVITY

<u>HIV</u>	<u>VACCINIA</u>	<u>PUNTA TORO</u>	<u>YELLOW FEVER</u>
6460	6462	6422	6456
6457	6467	6443	6458
6455		6444	6444
6466		6445	6445
		6402	

APPENDIX VI

SUMMARY OF SIGNIFICANT PROJECT ACCOMPLISHMENTS TO DATE AND RECOMMENDATIONS FOR FUTURE EXPLOITATION OF FINDINGS

PROJECT TITLE: SUICIDE INHIBITORS OF VIRAL POLYMERASES AS VIRAL PROPHYLACTICS AND BIOLOGICAL WARFARE ANTIDOTES.

a) Problems to be studied. This project will synthesize new types of antiviral drugs based upon suicide inhibitors of viral polymerases. These compounds will be screened in collaborative studies with the U.S. Army Antiviral Testing Facility for antiviral activity against a spectrum of 10 viruses of interest as military disease hazards and biological warfare agents.

b) Significance and Uniqueness. Suicide and affinity inhibitors of both DNA and RNA viral polymerases will be synthesized. This type of inhibitor contains a latent reactive moiety which selectively and irreversibly inactivates the viral enzyme. In preliminary studies, about 30 compounds with these potentialities have been synthesized and screened for antiviral activity in tissue culture. A number of active compounds including a new family - the nucleoside spiroxiranes - have been identified. Cytotoxicity assays in cultured T-lymphocytes also indicate favorable therapeutic indices for these types of drugs.

Two compounds, 2'3' sulfanyl cytidine hydrochloride and 2',0² anhydrocytidine hydrochloride, which have proved to be highly effective against vaccinia virus in tissue culture, will be synthesized in larger quantities for further characterization and *in vivo* studies in the U.S. Army Antiviral Facility, Fort Detrick, MD. Congeners of compounds that have shown moderate activity against Punta Toro and yellow fever viruses will also be developed. Test data on selected compounds are given in the appendix.

Preliminary studies have begun on a series of nucleoside 5' oxaphosphorins and dioxaphospholes that are suicide analogs directed against enzymatic displacement reactions at the 5' α phosphate of the nucleotide substrate. The action mechanism and selectivity of the drugs will be characterized against viral and host nucleotide polymerases *in vitro*.

In the continuing studies, the range and selectivity of the nucleoside spiroxiranes will be extended by synthesizing additional members of the family. The suicide nature of their action and sensitivity will be determined in kinetic studies using viral and cellular DNA and RNA polymerases *in vitro*. Samples (75 mg) of each compound will be supplied to USAMRID for *in vitro* testing. Larger samples (2 g) of compounds showing activity *in vitro* will be supplied for further testing *in vivo*.

c) Relevance to USAMRDC mission studies. Suicide inhibitors represent a new class of antiviral drugs potentially capable of great selectivity. Orally administered antivirals could be militarily useful for temporary viral prophylaxis in emergency troop deployments to environments where advance vaccination is not possible, as antidotes following battlefield exposure to vaccinia-based or other biological warfare vectors, and in other unanticipated epidemic situations.

d) Estimated Project Duration and Personnel. Organic chemist (50% time), 1 or 2 graduate students. Duration 3 years.

e) Animal use. No animal or human use except USAMRDC in-house testing of antivirals supplied as requested.